385 390 395 400

Arg Asn Asn Leu Asn Asn Arg Val Asn Ser Gly Ser Ser Ser Asn Ile 405 410 415

Ser Asn Thr Ala Ala Asn His Pro Tyr Gly Ala Pro Glu Phe Asn Met 420 425 430

Ile Ala Asn Asn Thr Pro Ala Ala Leu Thr Tyr Asn Arg Ala His Phe
435 440 445

Pro Ala Ile Thr Pro Leu Ser Arg Gln Asn Ser Leu Asn Met Ala Pro 450 455 460

Ser Asn Ser Gly Ser Pro Ile Ile Ile Ala Asp His Phe Ser Gly Asn 465 470 475 480

Asn Asn Ile Ala Pro Asn Tyr Arg Tyr Asn Asn Asn Ile Asn Asn Asn 485 490 495

Asn Asn Asn Ile Asn Asn Met Thr Asn Asn Arg Tyr Asn Ile Asn Asn Asn 500 505 510

Asn Ile Asn Gly Asn Gly Asn Gly Asn Gly Asn Asn Ser Asn Asn Asn 515 520 525

Asn Asn His Asn Asn Asn His Asn Asn His His Asn Gly Ser Ile 530 535 540

Asn Ser Asn Ser Asn Thr Asn Asn Asn Asn Asn Asn Asn Asn Gly Asn 545 550 560

Asn Ser Asn Asn Cys Asn Ser Asn Ile Gly Met Gly Gly Cys Gly Ser 565 570 575

Asn Met Pro Phe Arg Ala Gly Asp Trp Lys Cys Ser Thr Cys Thr Tyr 580 585 590

His Asn Phe Ala Lys Asn Val Val Cys Leu Arg Cys Gly Gly Pro Lys 595 600 605

Ser Ile Ser Gly Asp Ala Ser Glu Thr Asn His Tyr Ile Asp Ser Ser 610 615 620

Thr Phe Gly Pro Ala Ser Arg Thr Pro Ser Asn Asn Asn Ile Ser Val 625 630 635 640

Asn Thr Asn Gly Gly Ser Asn Ala Gly Arg Thr Asp Gly Asn Asp Asn

645

650

655

Lys Gly Arg Asp Ile Ser Leu Met Glu Phe Met Ser Pro Pro Leu Ser 660 665 670

Met Ala Thr Lys Ser Met Lys Glu Gly Asp Gly Asn Gly Ser Ser Phe 675 680 685

Asn Glu Phe Lys Ser Asp Lys Ala Asn Val Asn Phe Ser Asn Val Gly 690 695 700

Asp Asn Ser Ala Phe Gly Asn Gly Phe Asn Ser Ser Ile Arg Trp
705 710 715

<210> 47

<211> 578

<212> DNA

<213> Saccharomyces cerevisiae

<400> 47

aacaagaaaa cccttccgtt gatcttagat ttcttagagg ttcatgaact agaaagcgac 60 ttgaacaaag tcatccttca agaatcgaaa tctcaagaga aaaacaaaat ttaatgtaga 120 ttgtccacta tctcatgtaa atatacataa caggtattcc tgagcgttcg aaaattagac 180 tgtacttttc tgatgcgctc ccgtacacct ttgacatata caaacatccg cacattttat 240 agctttcttt ctagaatttt ttccacgcgc tctcgatcaa tgaactctta aaagtaactg 300 acccctaact ttttcccagg caaggctggc ctcattaccc tacccgaaag tttcacttta 360 ccccatggca gatggacgat attttaacaa gacgaaactg acctcgtct ctataaaact 420 ggacttctaa gcaactctca tttatcttat atccgttcca ttttgtacta aaagaaccag 480 accacatcga ttcaatcgaa atgagagca agtggagaaa gaagagaact agaagactta 540 agagaaaagag acggaaggtg agagccagat ccaaataa

<210> 48

<211> 25

<212> PRT

<213> Saccharomyces cerevisiae

<400> 48

Met Arg Ala Lys Trp Arg Lys Lys Arg Thr Arg Arg Leu Lys Arg Lys

1 10 15

Arg Arg Lys Val Arg Ala Arg Ser Lys
20 25

```
<211> 1354
<212> DNA
<213> Saccharomyces cerevisiae
<400> 49
tattgacgtt tcgctctcag gtccaccgtg ttctcaaaag atacttttaa aacctaaaac 60
acacgaaatc atattatgat aattcagaat gatagtgtgg tactgtgtca attgactgtt 120
caagactgaa gaggatettt gatttgttgt tactcaacaa ataatettea egaaaaettt 180
ctcaatctgg ggactgtatt aatctcagac ccatacatat ctacacccat aactttttac 240
atttaatttt ttatcacata ataggtagct taaattgtaa agtcgcaaaa aaaaatggca 300
gcgcagcctc tccgggtgaa ccccacgaca acttacctgg cactccatgc actaacgggc 360
gggtttgggc aggattccag catcaatttt gcaaaattca cacctgagta attcatatat 420
gtaatataat gttaagcata cgctgtcgat tagcactatt attgaccgta gaataggtac 480
aqtqaqacaq tatattcgaa atggtatgtt tgagatgaac aaaataataa agactgacaa 540
ctgcagaaca gagaggatca tagcaaccta gtgcaagaaa gagcctcgaa gcgttaaact 600
tttggaacgt aattcatccg ttggctatgt ctattcaata cagttagaat acgaaagctg 660
taatcaagta tatcggatta ttcgcaagca aagaatcaag gaaaagaaag tgaaaatagc 720
atacatcttt aaattcagag gttttgctga attttaatag ggaagtttac gttatgattg 780
ttggccgtag atcgcctagc gtttgaccaa atttaattct cctaattttt ttattgtaaa 840
aaagccttcc aacgaaataa attagttatt ggtttttttt ctctgttacg agggatatat 900
gatgcctgtg cttgttagtt cattataagt gctaataaaa tactaacgtt aataaaatt 960
tggaatatta tttcattttt tatcctatta ataggccggt gttaaagctt acgaactaag 1020
aaccaaatcc aaggaacaat tggcttctca attggttgac ttgaaaaagg agttggctga 1080
attgaaggtc caaaagttgt ccagaccatc tttgccaaag atcaagaccg tcagaaagag 1140
tatcgcctgt gtcttgaccg tcatcaacga acaacaaaga gaagctgtta gacaattata 1200
caagggtaag aagtaccaac caaaggactt gagagccaag aagaccagag ctttgagaag 1260
agetttgace aaattegaag etteecaagt taeegaaaag caaagaaaga ageaaatege 1320
                                                                   1354
tttcccacaa agaaagtacg ctattaaggc ttaa
<210> 50
<211> 120
<212> PRT
<213> Saccharomyces cerevisiae
<400> 50
Met Ala Gly Val Lys Ala Tyr Glu Leu Arg Thr Lys Ser Lys Glu Gln
                  5
  1
Leu Ala Ser Gln Leu Val Asp Leu Lys Lys Glu Leu Ala Glu Leu Lys
                                 25
                                                     30
             20
Val Gln Lys Leu Ser Arg Pro Ser Leu Pro Lys Ile Lys Thr Val Arg
        35
                             40
                                                 45
Lys Ser Ile Ala Cys Val Leu Thr Val Ile Asn Glu Gln Gln Arg Glu
     50
                         55
                                             60
```

02-07-1999 (EP998/0141.1)

Ala Val Arg Gln Leu Tyr Lys Gly Lys Lys Tyr Gln Pro Lys Asp Leu 65 70 75 80

Arg Ala Lys Lys Thr Arg Ala Leu Arg Arg Ala Leu Thr Lys Phe Glu 85 90 95

Ala Ser Gln Val Thr Glu Lys Gln Arg Lys Lys Gln Ile Ala Phe Pro 100 105 110

Gln Arg Lys Tyr Ala Ile Lys Ala 115 120

<210> 51

<211> 3254

<212> DNA

<213> Saccharomyces cerevisiae

<400> 51

atcaagtttc ctttaaaggg atatataaca gattctaaaa ctgacagaaa tatttcgagt 60 gaagaagaag cgttaaatat tggatctttc cgcagttcta ctctgataca tttttgaagt 120 aggagagtca tttagaaggc gtattgctca atagtagaaa gcaggcctgt gcacatgaat 180 taattaaaaa atataaaggt agtgattaga cgacacatgt ccataggtaa cctgtcataa 240 ttttgaacaa tttcccttct tttcttttt ttttttgggt gcggcgatat gtagcttgtt 300 aatttacaca tcatgtactt ttctgcatca aaatatgaaa ggcgatagta gctaaagaaa 360 ataccgagaa tttcctcgaa aagttgacga caaaagaaag gcataaaaaa gtaatttgaa 420 aatattttaa aactgtttta acccatctag catccgcgct aaaaaaggaa gatacaggat 480 acageggaaa caacttttaa atgatggaaa eteetacaga caatatagtt teeeetttte 540 acaattttgg tagctcgaca caatatagtg gtaccttgtc gagaactccc aaccaaataa 600 tagagctaga gaagcccagt actctatccc cattgtcaag aggaaaaaaa tggacggaaa 660 cctcctctac attttcgttc tcacccaaat ctagggtcac ttcttcaaac tcttctggca 780 atgaagacgg taacctaatg aatacacctt ctacggtttc cactgattat ttgccacaac 840 acceteacag aacategtet ttgecaagae etaattecaa tetettteae geaagtaata 900 gtaacctatc ccgagcaaat gagcccccaa gggccgaaaa tttatcagat aatataccac 960 ccaaggtcgc tccatttggc tatccaatac aaagaacctc tattaaaaaa tcctttttga 1020 atgettettg taegttatgt gaegageeta tttetaacag aagaaaggga gagaaaatta 1080 tagagettge atgtggeeae ttaagteaee aagaatgtet tattatetet ttttggeaeea 1140 cttcaaaggc agacgttcgt gcgctatttc ctttttgtac caaatgtaaa aaagatacta 1200 acaaagccgt tcaatgcatt ccagaaaatg atgaactaaa ggatattcta atttctgatt 1260 ttttgattca taagattcct gattctgagt tatcaatcac acctcagtcc cgctttcctc 1320 cttattcacc actcttgcct ccttttgggt tatcctatac acctgttgaa agacaaacga 1380 tatattetea agetecaagt etaaaceeaa ateteatatt ggetgeacee eecaaggaaa 1440 gaaaccaaat tccacaaaaa aaatcaaact atacattttt acattcaccc ctggggcaca 1500 gaagaattcc gtccggagca aactctatct tagcagacac ctctgtagcg ttgtcagcta 1560 atgattetat ttetgetgtt tecaattegg taagageaaa ggatgaegaa aecaaaacaa 1620

```
cgttgccgct gttaaggtca tattttattc aaattctttt gaacaatttc caggaagaat 1680
tgcaggattg gagaatagac ggggactatg gattactaag gttggtagac aaattgatga 1740
tttccaaaga tggtcagaga tatatacaat gctggtgttt cttatttgaa gacgcatttg 1800
taatagcaga agtggataac gatgttgatg ttttggaaat tagactaaag aatttagaag 1860
tatttacacc tattgccaac ttgagaatga ctacactcga agettcagta ctcaaatgca 1920
ccttaaataa acaacattgc gccgatttat cagatcttta cattgttcag aatataaatt 1980
ctgacgaaag cacaactgta cagaaatgga tatcaggtat attgaatcag gattttgtat 2040
tcaatgagga caatatcact tcgaccctgc ctattcttcc cattataaag aacttttcaa 2100
aagatgttgg taatggtagg cacgagacga gtacctttct aggtttaatc aatcctaaca 2160
aagttgttga agttggaaat gtgcacgata atgatactgt aatcataagg aggggattca 2220
ccttaaattc aggagaatgt tctaggcaga gtactgtcga cagtatacaa tctgttctaa 2280
ccacgataag ctcaattctt tcccttaaac gagaaaaacc tgataatttg gcaataatct 2340
tacagatcga ttttacgaaa ttgaaggaag aagacagttt aattgttgtt tataacagtc 2400
taaaagcttt aaccattaaa tttgcgcgtt tgcagttttg tttcgttgat cgaaataatt 2460
atgttctgga ctatggatcg gtattacaca agatagattc actagattcc atctcaaatc 2520
tcaaatcaaa gagttcctcg acacaatttt cacctatttg gttgaaaaat actctatatc 2580
ccgaaaatat tcatgaacat ttgggtattg ttgctgtatc aaatagtaat atggaagcaa 2640
aaaaatccat actatttcaa gattacagat gctttacaag ttttggaaga agaaggccca 2700
atgaattgaa gattaaggtg ggctatttga acgttgacta cagtgataaa attgatgaac 2760
tagtegagge cagetectgg actititgtit tagaaactet tigetacagt tieggtetaa 2820
gttttgatga acatgatgac gatgacgaag aggataatga tgattcgacc gataatgaac 2880
ttgataatag ttcaggatca ctgtcggatg ctgaatctac aactactatt catattgatt 2940
ctccatttga taatgaaaat gctaccgcaa atatggtgaa tgacagaaac cttctcactg 3000
agggtgaaca tagcaatata gaaaacttag aaactgtcgc ttcttcagta cagccagctc 3060
tgattcctaa tattagattt tcacttcatt ctgaggagga aggtactaat gaaaatgaaa 3120
atgaaaatga tatgccagta ttattactta gtgatatgga taaaggaatc gatggcataa 3180
ccagacgcag ttcattctcg agtcttatag agagcggtaa taacaactgt cccctccata 3240
                                                                  3254
tggattatat atag
```

```
<210> 52
```

<211> 917

<212> PRT

<213> Saccharomyces cerevisiae

## <400> 52

Met Met Glu Thr Pro Thr Asp Asn Ile Val Ser Pro Phe His Asn Phe 1 5 10 15

Gly Ser Ser Thr Gln Tyr Ser Gly Thr Leu Ser Arg Thr Pro Asn Gln
20 25 30

Ile Ile Glu Leu Glu Lys Pro Ser Thr Leu Ser Pro Leu Ser Arg Gly
35 40 45

Lys Lys Trp Thr Glu Lys Leu Ala Arg Phe Gln Arg Ser Ser Ala Lys 50 55 60



<b>L</b> ys 65	Lys	Arg	Phe	Ser	Pro 70	Ser	Pro	Ile	Ser	Ser 75	Ser	Thr	Phe	Ser	Phe 80
Ser	Pro	Lys	Ser	Arg 85	Val	Thr	Ser	Ser	Asn 90	Ser	Ser	Gly	Asn	Glu 95	Asp
Gly	Asn	Leu	Met 100	Asn	Thr	Pro	Ser	Thr 105	Val	Ser	Thr	Asp	Туг 110	Leu	Pro
Gln	His	Pro 115	His	Arg	Thr	Ser	Ser 120	Leu	Pro	Arg	Pro	Asn 125	Ser	Asn	Leu
Phe	His 130	Ala	Ser	Asn	Ser	Asn 135	Leu	Ser	Arg	Ala	Asn 140	Glu	Pro	Pro	Arg
Ala 145	Glu	Asn	Leu	Ser	Asp 150	Asn	Ile	Pro	Pro	Lys 155	Val	Ala	Pro	Phe	Gly 160
Tyr	Pro	Ile	Gln	Arg 165	Thr	Ser	Ile	Lys	Lys 170	Ser	Phe	Leu	Asn	Ala 175	Ser
Cys	Thr	Leu	Суs 180	Asp	Glu	Pro	Ile	Ser 185	Asn	Arg	Arg	Lys	Gly 190	Glu	Lys
Ile	Ile	Glu 195	Leu	Ala	Суѕ	Gly	His 200	Leu	Ser	His	Gln	Glu 205	Суѕ	Leu	Ile
Ile	Ser 210		Gly	Thr	Thr	Ser 215	Lys	Ala	Asp	Val	Arg 220	Ala	Leu	Phe	Pro
Phe 225	Суѕ	Thr	Lys	Суѕ	Lys 230	Lys	Asp	Thr	Asn	Lys 235	Ala	Val	Gln	Cys	11e 240
			Asp	245		_	_		250			_		255	
			Pro 260					265					270		
		275	Ser				280					285	_		
	290		Gln			295					300				
Leu	Ile	Leu	Ala	Ala	Pro	Pro	Lys	Glu	Arg	Asn	Gln	Ile	Pro	Gln	Lys

Lys	Ser	Asn	Tyr	Thr	Phe	Leu	His	Ser	Pro	Leu	Gly	His	Arg	Arg 335	Ile
-				325					330					333	

- Pro Ser Gly Ala Asn Ser Ile Leu Ala Asp Thr Ser Val Ala Leu Ser 340 345 350
- Ala Asn Asp Ser Ile Ser Ala Val Ser Asn Ser Val Arg Ala Lys Asp 355 360 365
- Asp Glu Thr Lys Thr Thr Leu Pro Leu Leu Arg Ser Tyr Phe Ile Gln 370 375 380
- Ile Leu Leu Asn Asn Phe Gln Glu Glu Leu Gln Asp Trp Arg Ile Asp 385 390 395 400
- Gly Asp Tyr Gly Leu Leu Arg Leu Val Asp Lys Leu Met Ile Ser Lys
  405 410 415
- Asp Gly Gln Arg Tyr Ile Gln Cys Trp Cys Phe Leu Phe Glu Asp Ala 420 425 430
- Phe Val Ile Ala Glu Val Asp Asn Asp Val Asp Val Leu Glu Ile Arg 435 440 445
- Leu Lys Asn Leu Glu Val Phe Thr Pro Ile Ala Asn Leu Arg Met Thr 450 455 460
- Thr Leu Glu Ala Ser Val Leu Lys Cys Thr Leu Asn Lys Gln His Cys 465 470 475 480
- Ala Asp Leu Ser Asp Leu Tyr Ile Val Gln Asn Ile Asn Ser Asp Glu
  485 490 495
- Ser Thr Thr Val Gln Lys Trp Ile Ser Gly Ile Leu Asn Gln Asp Phe 500 505 510
- Val Phe Asn Glu Asp Asn Ile Thr Ser Thr Leu Pro Ile Leu Pro Ile 515 520 525
- Ile Lys Asn Phe Ser Lys Asp Val Gly Asn Gly Arg His Glu Thr Ser 530 535
- Thr Phe Leu Gly Leu Ile Asn Pro Asn Lys Val Val Glu Val Gly Asn 545 550 555 560
- Val His Asp Asn Asp Thr Val Ile Ile Arg Arg Gly Phe Thr Leu Asn 565 570 575



Ser	Gly	Glu	Суs 580	Ser	Arg	Gln	Ser	Thr 585	Val	Asp	Ser	Ile	Gln 590	Ser	Val
Leu	Thr	Thr 595	Ile	Ser	Ser	Ile	Leu 600	Ser	Leu	Lys	Arg	Glu 605	Lys	Pro	Asp
Asn	Leu 610	Ala	Ile	Ile	Leu	Gln 615	Ile	Asp	Phe	Thr	Lys 620	Leu	Lys	Glu	Glu
Asp 625	Ser	Leu	Ile	Val	Val 630	Tyr	Asn	Ser	Leu	Lys 635	Ala	Leu	Thr	Ile	Lys 640
Phe	Ala	Arg	Leu	Gln 645	Phe	Суз	Phe	Val	Asp 650	Arg	Asn	Asn	Tyr	Val 655	Leu
Asp	Tyr	Gly	Ser 660	Val	Leu	His	Lys	Ile 665	Asp	Ser	Leu	Asp	Ser 670	Ile	Ser
Asn	Leu	Lys 675	Ser	Lys	Ser	Ser	Ser 680	Thr	Gln	Phe	Ser	Pro 685	Ile	Trp	Leu
Lys	Asn 690	Thr	Leu	Tyr	Pro	G1u 695	Asn	Ile	His	Glu	His 700	Leu	Gly	Ile	Val
Ala 705	Val	Ser	Asn	Ser	Asn 710	Met	Glu	Ala	Lys	Lys 715	Ser	Ile	Leu	Phe	Gln 720
Asp	Tyr	Arg	Cys	Phe 725	Thr	Ser	Phe	Gly	Arg 730	Arg	Arg	Pro	Asn	Glu 735	Leu
Lys	Ile	Lys	Val 740	Gly	Tyr	Leu	Asn	Val 745	qaA	Tyr	Ser	Asp	Lys 750	Ile	Asp
Glu	Leu	Val 755	Glu	Ala	Ser	Ser	Trp 760	Thr	Phe	Val	Leu	Glu 765	Thr	Leu	Cys
Tyr	Ser 770	Phe	Gly	Leu	Ser	Phe 775	Asp	Glu	His	Asp	Asp 780	Asp	Asp	Glu	Glu
Asp 785	Asn	Asp	Asp	Ser	Thr 790	Asp	Asn	Glu	Leu	Asp 795	Asn	Ser	Ser	Gly	Ser 800
Leu	Ser	Asp	Ala	Glu	Ser	Thr	Thr	Thr	Ile	His	Ile	Asp	Ser	Pro	Phe

Asp Asn Glu Asn Ala Thr Ala Asn Met Val Asn Asp Arg Asn Leu Leu

```
Thr Glu Gly Glu His Ser Asn Ile Glu Asn Leu Glu Thr Val Ala Ser
        835
                            840
                                                 845
Ser Val Gln Pro Ala Leu Ile Pro Asn Ile Arg Phe Ser Leu His Ser
    850
                        855
                                             860
Glu Glu Glu Gly Thr Asn Glu Asn Glu Asn Glu Asn Asp Met Pro Val
                    870
                                         875
Leu Leu Leu Ser Asp Met Asp Lys Gly Ile Asp Gly Ile Thr Arg Arg
                885
                                     890
Ser Ser Phe Ser Ser Leu Ile Glu Ser Gly Asn Asn Asn Cys Pro Leu
                                905
His Met Asp Tyr Ile
        915
```

```
<210> 53
<211> 3422
<212> DNA
<213> Saccharomyces cerevisiae
```

<400> 53

```
ccgtgtcaag atctaacacg gtagtcagct actacacaag gtctcagaac agaatgagaa 60
gtggaacact ggataatgat tacgtgaaca gacaaaagct tcctacacat atctctcttc 120
aagattatcg tgatgctaat gctagaagta atatatcgcg tcaggactct gtctccacaa 180
cgaactctga tgtggtagac ctcagctatt ctctggggca tggcttgcgt gtggcaaacc 240
ctgattcaga cccagaatga tatccttctc tgtagttttg tagatgtcat atatgtacgt 300
ttatggaaca gcattttaga aaagtattac ccagcttatc acttcgtttt ttttttcttt 360
ccgtgtactc gctacacgta gaaagaaatc agaaaacaac agctcgacaa gtgaaatttg 420
acgttcatta agactcagtt aagattgcct tgagaataaa caaaagtaat cacagttaac 480
tattgaacaa gagtgcacct atgacttcac tttcttcaca gccagcgtac acgttggttt 540
tcgatccttc tccgagtatg gagacttact cgagtaccga ttttcagaaa gctcttgaaa 600
agggatctga tgaacaaaaa attgacacga tgaaatcaat tttagttaca atgctggaag 660
gaaatccaat gcctgaattg ttgatgcaca taataagatt tgtcatgcct tctaaaaata 720
aggaattaaa aaagcttttg tacttctact gggaaattgt tcccaaacta gctgaagatg 780
gaaaattgag acatgaaatg attcttgtct gtaatgccat tcaacacgat ttgcaacatc 840
ctaatgaata tattagaggt aacacattaa ggtttttaac gaaattgaga gaggccgaac 900
tcttagaaca gatggttccc tctgtcttag cgtgcttgga ataccgtcat gcatatgttc 960
gtaagtatgc aatcetagca gttttctcca ttttcaaggt cagcgaacat ttacttcccg 1020
atgctaaaga aatcatcaat tcgttcatag tagctgaaac tgatccaata tgtaaaagaa 1080
atgcatttat tgggttagct gaattagatc gtgaaaatgc cttacactat ttagagaaca 1140
atattgctga tatagaaaac ctagaccctt tattacaagc tgtctttgtt caatttatca 1200
gacaagatgc aaacaggacc cctgctttga aagcccaata tatcgaatta ttgatggaac 1260
tgctttcgac cacgacttcc gatgaagtca tcttcgagac cgcattagcc ctaactgtgt 1320
```

```
tgtctgccaa tccaaatgtc ttggttcctg cggttaacaa attgattgac ttggccgtca 1380
aggtttctga taataacatt aagttaattg ttctagaccg tattcaagac atcaatgcta 1440
ataacgtagg tgctttggaa gagttaaccc tggatatttt gagagtcttg aatgcagaag 1500
atttagacgt tcgttcaaag gcgcttgata tttcaatgga cttggccaca tccagaaatg 1560
ctgaagatgt tgttcagctt ttgaagaaag agctgcaaac aaccgtaaat aacccagatc 1620
aagacaaggc aatgcagtac agacaattgt taataaaaac tattcgtacc gtggctgtaa 1680
actttgtaga aatggcagca agtgttgttt cgctattatt agatttcatc ggtgatttaa 1740
acteggttge egecagtggt atcattgeet ttateaaaga agtgategaa aaataeccae 1800
aacttagagc caatateett gaaaacatgg tteaaaeget agacaaagtg agatetgeta 1860
aagettaceg eggtgeatta tggattatgg gtgagtatge tgaaggagaa agtgagatac 1920
aacattgttg gaagcacatt cgtaacagcg taggtgaagt teetateett caatcagaaa 1980
tcaaaaagtt aacacaaaac caagaacaca ccgaagaaaa tgaggttgac gctaccgcca 2040
agccaactgg tccagttatt ctaccagacg gtacgtatgc cactgaaagc gctttcgatg 2100
tgaagacttc tcaaaagtca gttaccgatg aagaacgtga ttctagacct ccaattcgcc 2160
ggtttgtttt aagtggtgat ttctacacag ctgccattct ggccaacacc atcattaaac 2220
ttgttttaaa attcgaaaac gtttccaaga acaaaactgt catcaatgct ctaaaggcgg 2280
aagctttact aattttagtt agtattgtaa gagtgggtca aagctctttg gtggagaaaa 2340
aaattgatga agattettta gagagagtta tgacatetat ttetatttta ttggatgaag 2400
ttaatcctga ggaaaagaag gaagaagtta aacttctgga ggttgcattc ctggacacca 2460
ccaaatcctc attcaagaga caaattgaaa ttgcaaagaa gaacaagcat aagagagcat 2520
taaaagacag ttgcaaaaac atcgaaccaa ttgatacgcc gatttctttc aggcaatttg 2580
ctggtgtgga ttctactaat gtgcaaaaag atagtattga agaagattta caactggcaa 2640
tgaaaggaga tgcaatccac gctactagca gctctagtat ttcgaagctg aagaagatag 2700
tacctttatg tggcttttct gatccagttt acgccgaggc ttgtattaca aacaatcaat 2760
ttgacgtcgt attagatgtt cttcttgtta atcaaacgaa agaaacattg aaaaacctac 2820
atgtgcaatt tgcaactett ggtgatttga agattattga cacaccacag aagaccaacg 2880
tgattcctca tggcttccac aaattcactg ttactgtcaa agtttcctct gctgacacag 2940
gtgtcatttt cggtaatatt atttatgatg gtgcgcatgg tgaagatgct cgttatgtta 3000
ttttaaacga cgttcatgtt gacattatgg attatatcaa accagccact gctgacgatg 3060
aacattteeg taccatgtgg aatgeatttg agtgggagaa caaaatateg gtcaaatcac 3120
aactaccaac attgcatgct tatttgagag aactggtcaa gggaactaat atgggtattc 3180
taacaccatc agagtcgttg ggagaagatg attgtaggtt cttaagttgt aatctgtatg 3240
cgaagtcgtc ctttggtgaa gatgccctag ccaacttgtg tatcgaaaag gattccaaaa 3300
ccaatgatgt cataggttat gttcgtatcc gatcaaaggg acaaggtttg gctctgtccc 3360
taggtgacag agtggcattg attgctaaga agaccaataa acttgctctc actcatgttt 3420
                                                                  3422
ga
```

```
<210> 54 <211> 973
```

<212> PRT

<213> Saccharomyces cerevisiae

<400> 54

Met Thr Ser Leu Ser Ser Gln Pro Ala Tyr Thr Leu Val Phe Asp Pro 1 5 10 15

Ser Pro Ser Met Glu Thr Tyr Ser Ser Thr Asp Phe Gln Lys Ala Leu

20

25

- Glu Lys Gly Ser Asp Glu Gln Lys Ile Asp Thr Met Lys Ser Ile Leu 35 40 45
- Val Thr Met Leu Glu Gly Asn Pro Met Pro Glu Leu Leu Met His Ile
  50 55 60
- Ile Arg Phe Val Met Pro Ser Lys Asn Lys Glu Leu Lys Lys Leu Leu 65 70 75 80
- Tyr Phe Tyr Trp Glu Ile Val Pro Lys Leu Ala Glu Asp Gly Lys Leu
  85 90 95
- Arg His Glu Met Ile Leu Val Cys Asn Ala Ile Gln His Asp Leu Gln
  100 105 110
- His Pro Asn Glu Tyr Ile Arg Gly Asn Thr Leu Arg Phe Leu Thr Lys
  115 120 125
- Leu Arg Glu Ala Glu Leu Leu Glu Gln Met Val Pro Ser Val Leu Ala 130 135 140
- Cys Leu Glu Tyr Arg His Ala Tyr Val Arg Lys Tyr Ala Ile Leu Ala 145 150 155 160
- Val Phe Ser Ile Phe Lys Val Ser Glu His Leu Leu Pro Asp Ala Lys 165 170 175
- Glu Ile Ile Asn Ser Phe Ile Val Ala Glu Thr Asp Pro Ile Cys Lys 180 185 190
- Arg Asn Ala Phe Ile Gly Leu Ala Glu Leu Asp Arg Glu Asn Ala Leu 195 200 205
- His Tyr Leu Glu Asn Asn Ile Ala Asp Ile Glu Asn Leu Asp Pro Leu 210 215 220
- Leu Gln Ala Val Phe Val Gln Phe Ile Arg Gln Asp Ala Asn Arg Thr 225 230 235 240
- Pro Ala Leu Lys Ala Gln Tyr Ile Glu Leu Leu Met Glu Leu Leu Ser 245 250 255
- Thr Thr Thr Ser Asp Glu Val Ile Phe Glu Thr Ala Leu Ala Leu Thr 260 265 270
- Val Leu Ser Ala Asn Pro Asn Val Leu Val Pro Ala Val Asn Lys Leu

275



- Ile Asp Leu Ala Val Lys Val Ser Asp Asn Asn Ile Lys Leu Ile Val 290 295 300
- Leu Asp Arg Ile Gln Asp Ile Asn Ala Asn Asn Val Gly Ala Leu Glu 305 310 315 320
- Glu Leu Thr Leu Asp Ile Leu Arg Val Leu Asn Ala Glu Asp Leu Asp 325 330 335
- Val Arg Ser Lys Ala Leu Asp Ile Ser Met Asp Leu Ala Thr Ser Arg 340 345 350
- Asn Ala Glu Asp Val Val Gln Leu Leu Lys Lys Glu Leu Gln Thr Thr 355 360 365
- Val Asn Asn Pro Asp Gln Asp Lys Ala Met Gln Tyr Arg Gln Leu Leu 370 375 380
- Ile Lys Thr Ile Arg Thr Val Ala Val Asn Phe Val Glu Met Ala Ala 385 390 395 400
- Ser Val Val Ser Leu Leu Leu Asp Phe Ile Gly Asp Leu Asn Ser Val 405 410 415
- Ala Ala Ser Gly Ile Ile Ala Phe Ile Lys Glu Val Ile Glu Lys Tyr 420 425 430
- Pro Gln Leu Arg Ala Asn Ile Leu Glu Asn Met Val Gln Thr Leu Asp 435 440 445
- Lys Val Arg Ser Ala Lys Ala Tyr Arg Gly Ala Leu Trp Ile Met Gly 450 455 460
- Glu Tyr Ala Glu Gly Glu Ser Glu Ile Gln His Cys Trp Lys His Ile 465 470 475 480
- Arg Asn Ser Val Gly Glu Val Pro Ile Leu Gln Ser Glu Ile Lys Lys 485 490 495
- Leu Thr Gln Asn Gln Glu His Thr Glu Glu Asn Glu Val Asp Ala Thr 500 505 510
- Ala Lys Pro Thr Gly Pro Val Ile Leu Pro Asp Gly Thr Tyr Ala Thr 515 520 525
- Glu Ser Ala Phe Asp Val Lys Thr Ser Gln Lys Ser Val Thr Asp Glu

SEQL

530

535

540

Glu Arg Asp Ser Arg Pro Pro Ile Arg Arg Phe Val Leu Ser Gly Asp 545 550 555 560

Phe Tyr Thr Ala Ala Ile Leu Ala Asn Thr Ile Ile Lys Leu Val Leu 565 570 575

Lys Phe Glu Asn Val Ser Lys Asn Lys Thr Val Ile Asn Ala Leu Lys 580 585 590

Ala Glu Ala Leu Leu Ile Leu Val Ser Ile Val Arg Val Gly Gln Ser 595 600 605

Ser Leu Val Glu Lys Lys Ile Asp Glu Asp Ser Leu Glu Arg Val Met 610 615 620

Thr Ser Ile Ser Ile Leu Leu Asp Glu Val Asn Pro Glu Glu Lys Lys 625 630 635 640

Glu Glu Val Lys Leu Leu Glu Val Ala Phe Leu Asp Thr Thr Lys Ser 645 650 655

Ser Phe Lys Arg Gln Ile Glu Ile Ala Lys Lys Asn Lys His Lys Arg 660 665 670

Ala Leu Lys Asp Ser Cys Lys Asn Ile Glu Pro Ile Asp Thr Pro Ile 675 680 685

Ser Phe Arg Gln Phe Ala Gly Val Asp Ser Thr Asn Val Gln Lys Asp 690 695 700

Ser Ile Glu Glu Asp Leu Gln Leu Ala Met Lys Gly Asp Ala Ile His 705 710 715 720

Ala Thr Ser Ser Ser Ser Ile Ser Lys Leu Lys Lys Ile Val Pro Leu
725 730 735

Cys Gly Phe Ser Asp Pro Val Tyr Ala Glu Ala Cys Ile Thr Asn Asn 740 745 750

Gln Phe Asp Val Val Leu Asp Val Leu Leu Val Asn Gln Thr Lys Glu 755 760 765

Thr Leu Lys Asn Leu His Val Gln Phe Ala Thr Leu Gly Asp Leu Lys 770 780

Ile Ile Asp Thr Pro Gln Lys Thr Asn Val Ile Pro His Gly Phe His

785

9870141:1] | SEQL

800

Lys Phe Thr Val Thr Val Lys Val Ser Ser Ala Asp Thr Gly Val Ile 805 810 815

795

Phe Gly Asn Ile Ile Tyr Asp Gly Ala His Gly Glu Asp Ala Arg Tyr 820 825 830

Val Ile Leu Asn Asp Val His Val Asp Ile Met Asp Tyr Ile Lys Pro 835 840 845

Ala Thr Ala Asp Asp Glu His Phe Arg Thr Met Trp Asn Ala Phe Glu 850 855 860

Trp Glu Asn Lys Ile Ser Val Lys Ser Gln Leu Pro Thr Leu His Ala 865 870 875 880

Tyr Leu Arg Glu Leu Val Lys Gly Thr Asn Met Gly Ile Leu Thr Pro 885 890 895

Ser Glu Ser Leu Gly Glu Asp Asp Cys Arg Phe Leu Ser Cys Asn Leu 900 905 910

Tyr Ala Lys Ser Ser Phe Gly Glu Asp Ala Leu Ala Asn Leu Cys Ile 915 920 925

Glu Lys Asp Ser Lys Thr Asn Asp Val Ile Gly Tyr Val Arg Ile Arg 930 935 940

Ser Lys Gly Gln Gly Leu Ala Leu Ser Leu Gly Asp Arg Val Ala Leu 945 950 955 960

Ile Ala Lys Lys Thr Asn Lys Leu Ala Leu Thr His Val 965 970

<210> 55

<211> 1652

<212> DNA

<213> Saccharomyces cerevisiae

<400> 55

aaactttgtt caagattatg ctttcttgt aattttaaac atagtcttgc acttatttt 60 actgcatgtg gataaaagtt tttcgaatcg tttgcaggca aatgtaaact attactttt 120 aatgatttct gatactcttt gattccattc tgtcatactt ttttctgcat ttgaaacgct 180 aattaagtat ttcttggccc tgcttcttt ccttttgatt tctttttta ttctcaagtt 240 tttaatttct ggagaatctc ttttttgtt tattttgtta acacagttag tggagccttg 300

```
tagtatcgag agtagactat ctttggaaag caatgcgagt tgagagattg tggagtgtac 360
tacagaacca atattaaaca caatctttcc tcaaactgta acaccgagtt tttttcccca 420
ccaatcgtga atccgatagc atatactttt gtctagaaat ttcaataaac aacagaataa 480
cgaagagtgc taagggacaa atgcaaaacc ctccgttgat tcgtcccgat atgtataatc 540
agggaagcag ctcaatggct acttataatg cctctgagaa gaatctaaat gagcatcctt 600
ctccgcaaat tgcacagccc agcacgtccc aaaagttacc ttatagaata aatcctacaa 660
ccactaatgg ggacaccgac atatctgtta acagcaatcc tatccagcct cctttgccaa 720
acttgatgca tctatctggt ccgtctgact atagatcgat gcatcaaagt cctatacatc 780
catcttatat catccctccg cattcaaatg aaagaaaaca atcagcttct tacaacagac 840
ctcaaaatgc tcatgttagt attcaacctt ccgtggtatt cccccctaaa agttattcca 900
tatettatge acettateaa ataaateeee etttaeeaaa tggaetteeg aaceagagea 960
tatctttgaa taaggagtat attgcagagg agcaactatc aaccctccca tctcgcaata 1020
ccagtgttac tactgcacct cettettte aaaacagtgc tgataccgct aaaaattcag 1080
ctgataataa tgataataat gataatgtaa ccaaacctgt tcctgataaa gacacccaac 1140
tcataagtag ttcaggcaaa actttaagaa atactagaag agctgctcaa aatagaaccg 1200
ctcaaaaggc atttagacaa aggaaagaaa aatacatcaa gaatctcgaa caaaaatcaa 1260
agatatttga cgatttacta gcagaaaata ataacttcaa atcattaaac gattcattaa 1320
qaaatgacaa caacatttta atagctcagc atgaagctat aaggaatgca attactatgt 1380
taagaagtga gtatgatgtc ttatgtaacg aaaacaacat gttgaagaat gagaatagta 1440
taataaaaaa tgaacacaac atgtcaagaa atgaaaatga aaacctaaaa cttgagaata 1500
aacgcttcca cgctgaatat atacgaatga tcgaggatat tgaaaatact aaaagaaagg 1560
aacaagaaca acgagatgaa atagagcaac taaaaaaaaa aataagatcc ctggaggaaa 1620
                                                                  1652
tagtagggag acacteggat agtgccaegt aa
```

<211> 383

<212> PRT

<213> Saccharomyces cerevisiae

<400> 56

Met Gln Asn Pro Pro Leu Ile Arg Pro Asp Met Tyr Asn Gln Gly Ser
1 5 10 15

Ser Ser Met Ala Thr Tyr Asn Ala Ser Glu Lys Asn Leu Asn Glu His
20 25 30

Pro Ser Pro Gln Ile Ala Gln Pro Ser Thr Ser Gln Lys Leu Pro Tyr
35 40 45

Arg Ile Asn Pro Thr Thr Thr Asn Gly Asp Thr Asp Ile Ser Val Asn 50 55 60

Ser Asn Pro Ile Gln Pro Pro Leu Pro Asn Leu Met His Leu Ser Gly 65 70 75 80

Pro Ser Asp Tyr Arg Ser Met His Gln Ser Pro Ile His Pro Ser Tyr 85 90 95 02-07-1999

Ile	Ile	Pro	Pro 100	His	Ser	Asn	Glu	Arg 105	Lys	Gln	Ser	Ala	Ser 110	Туr	Asn
Arg	Pro	Gln 115	Asn	Ala	His	Val	Ser 120	Ile	Gln	Pro	Ser	Val 125	Val	Phe	Pro
Pro	Lys 130	Ser	Tyr	Ser	Ile	Ser 135	Tyr	Ala	Pro	Tyr	Gln 140	Ile	Asn	Pro	Pro
Leu 145	Pro	Asn	Gly	Leu	Pro 150	Asn	Gln	Ser	Ile	Ser 155	Leu	Asn	Lys	Glu	Tyr 160
Ile	Ala	Glu	Glu	Gln 165	Leu	Ser	Thr	Leu	Pro 170	Ser	Arg	Asn	Thr	Ser 175	Val
Thr	Thr	Ala	Pro 180	Pro	Ser	Phe	Gln	Asn 185	Ser	Ala	Asp	Thr	Ala 190	Lys	Asn
Ser	Ala	Asp 195	Asn	Asn	Asp	Asn	Asn 200	Asp	Asn	Val	Thr	Lys 205	Pro	Val	Pro
Asp	Lys 210	Asp	Thr	Gln	Leu	Ile 215	Ser	Ser	Ser	Gly	Lys 220	Thr	Leu	Arg	Asn
Thr 225	Arg	Arg	Ala	Ala	Gln 230	Asn	Arg	Thr	Ala	Gln 235	Lys	Ala	Phe	Arg	Gln 240
				245					250					Ile 255	
Asp	Asp	Leu	Leu 260	Ala	Glu	Asn	Asn	Asn 265	Phe	Lys	Ser	Leu	Asn 270	Asp	Ser
		275					280					285		Ile	
Asn	Ala 290	Ile	Thr	Met	Leu	Arg 295	Ser	Glu	Tyr	Asp	Val 300	Leu	Cys	Asn	Glu
305					310					315				His	320
Met	Ser	Arg	Asn	Glu 325	Asn	Glu	Asn	Leu	Lys 330	Leu	Glu	Asn	Lys	Arg 335	Phe

340

His Ala Glu Tyr Ile Arg Met Ile Glu Asp Ile Glu Asn Thr Lys Arg

02-07-1999 EP99870141.1



```
Lys Glu Gln Glu Gln Arg Asp Glu Ile Glu Gln Leu Lys Lys Lys Ile
355 360 365
```

Arg Ser Leu Glu Glu Ile Val Gly Arg His Ser Asp Ser Ala Thr 370 375 380

<210> 57 <211> 2270 <212> DNA <213> Saccharomyces cerevisiae

## <400> 57

ccgacagtac gacttaaaaa acaaaaacaa cgtccaggtg gaaaaagctg ccgcaaatgg 60 tacacggttt agatcaacaa gatccaatac ccctaattac acatgaatgt gacgtttcct 120 aaaagaagta gcctgtataa tatagagaga ttttatatac tttcttaatg aattagactg 180 tttctacaag tatttgacac tggaaaaaaa gagaaagtac atagagattg gccaaatatt 240 taaatctaca cagttgccta tegittateg cettattett cagaaacatt teatcaacta 300 ctctggtgca ttatttcatg ttagttacca tatataccgg ccggccgcat cggggttttt 360 tgattgaaaa aattggtata ttttcagtac acatataaat aaaaccctca atttgcctct 420 ccaaccgtta taactattcc agatcctctt taccgagcaa gtaggctagc ttctgtaaag 480 ggatttttcc atctaataca atgagtggag tatcaaataa aacagtatca attaatggtt 540 ggtatggcat gccaattcat ttactaaggg aagaaggcga ctttgcccag tttatgattc 600 taaccatcaa cgaattaaaa atagccatac atggttacct cagaaatacc ccatggtaca 660 acatgttgaa ggattatttg tttgtgatct tttgttacaa gctaataagt aatttttttt 720 atctgttgaa agtttatggg ccggtgaggt tagcagtgag aacatacgag catagttcca 780 gaagattgtt tcgttggtta ttggactcac catttttgag gggtaccgta gaaaaggaag 840 tcacaaaggt caaacaatcg atcgaagacg aactaattag atcggactct cagttaatga 900 atttcccaca gttgccatcc aatgggatac ctcaggatga tgttattgaa gagctaaata 960 aattgaacga cttgatacca catacccaat ggaaggaagg aaaggtctct ggtgccgttt 1020 accacggtgg tgatgatttg atccacttac aaacaatcgc atacgaaaaa tattgcgttg 1080 ccaatcaatt acatcccgat gtctttcctg ccgtacgtaa aatggaatcc gaagtggttt 1140 ctatggtttt aagaatgttt aatgcccctt ctgatacagg ttgtggtacc acaacttcag 1200 gtggtacaga atccttgctt ttagcatgtc tgagcgctaa aatgtatgcc cttcatcatc 1260 gtggaatcac cgaaccagaa ataattgctc ccgtaactgc acatgctggg tttgacaaag 1320 ctgcttatta ctttggcatg aagctacgcc acgtggagct agatccaacg acatatcaag 1380 tggacctggg aaaagtgaaa aaattcatca ataagaacac aattttactg gtcggttccg 1440 ctccaaactt tcctcatggt attgccgatg atattgaagg attgggtaaa atagcacaaa 1500 aatataaact teetttacae gtegacagtt gtetaggtte etttattgtt teatttatgg 1560 aaaaggctgg ttacaaaaat ctgccattac ttgactttag agtcccggga gtcacctcaa 1620 tatcatgtga cactcataaa tatggatttg caccaaaagg ctcgtcagtt ataatgtata 1680 gaaacagcga cttacgaatg catcagtatt acgtaaatcc tgcttggact ggcgggttat 1740 atggctctcc tacattagca gggtccaggc ctggtgctat tgtcgtaggt tgttgggcca 1800 ctatggtcaa catgggtgaa aatgggtaca ttgagtcgtg ccaagaaata gtcggtgcag 1860 caatgaagtt taaaaaatac atccaggaaa acattccaga cctgaatata atgggcaacc 1920 ctagatattc agtcatttca ttttcttcaa agaccttgaa catacacgaa ctatctgaca 1980

**-7**:

```
ggttgtccaa gaaaggctgg catttcaatg ccctacaaaa gccggttgca ctacacatgg 2040 ccttcacgag attgagcgct catgttgtgg atgagatctg cgacatttta cgtactaccg 2100 tgcaagagtt gaagagcgaa tcaaattcta aaccatccc agacggaact agcgctctat 2160 atggtgtcgc cgggagcgtt aaaactgctg gcgttgcaga caaattgatt gtgggattcc 2220 tagacgcatt atacaagttg ggtccaggag aggataccgc caccaagtag 2270
```

<211> 589

<212> PRT

<213> Saccharomyces cerevisiae

<400> 58

Met Ser Gly Val Ser Asn Lys Thr Val Ser Ile Asn Gly Trp Tyr Gly

1 5 10 15

Met Pro Ile His Leu Leu Arg Glu Glu Gly Asp Phe Ala Gln Phe Met 20 25 30

Ile Leu Thr Ile Asn Glu Leu Lys Ile Ala Ile His Gly Tyr Leu Arg
35 40 45

Asn Thr Pro Trp Tyr Asn Met Leu Lys Asp Tyr Leu Phe Val Ile Phe 50 55 60

Cys Tyr Lys Leu Ile Ser Asn Phe Phe Tyr Leu Leu Lys Val Tyr Gly
65 70 75 80

Pro Val Arg Leu Ala Val Arg Thr Tyr Glu His Ser Ser Arg Arg Leu 85 90 95

Phe Arg Trp Leu Leu Asp Ser Pro Phe Leu Arg Gly Thr Val Glu Lys
100 105 110

Glu Val Thr Lys Val Lys Gln Ser Ile Glu Asp Glu Leu Ile Arg Ser 115 120 125

Asp Ser Gln Leu Met Asn Phe Pro Gln Leu Pro Ser Asn Gly Ile Pro 130 135 140

Gln Asp Asp Val Ile Glu Glu Leu Asn Lys Leu Asn Asp Leu Ile Pro 145 150 155 160

His Thr Gln Trp Lys Glu Gly Lys Val Ser Gly Ala Val Tyr His Gly
165 170 175

Gly Asp Asp Leu Ile His Leu Gln Thr Ile Ala Tyr Glu Lys Tyr Cys 180 185 190 Val Ala Asn Gln Leu His Pro Asp Val Phe Pro Ala Val Arg Lys Met 195 200 205

Glu Ser Glu Val Val Ser Met Val Leu Arg Met Phe Asn Ala Pro Ser 210 215 220

Asp Thr Gly Cys Gly Thr Thr Thr Ser Gly Gly Thr Glu Ser Leu Leu 225 230 235 240

Leu Ala Cys Leu Ser Ala Lys Met Tyr Ala Leu His His Arg Gly Ile 245 250 255

Thr Glu Pro Glu Ile Ile Ala Pro Val Thr Ala His Ala Gly Phe Asp 260 265 270

Lys Ala Ala Tyr Tyr Phe Gly Met Lys Leu Arg His Val Glu Leu Asp 275 280 285

Pro Thr Thr Tyr Gln Val Asp Leu Gly Lys Val Lys Lys Phe Ile Asn 290 295 300

Lys Asn Thr Ile Leu Leu Val Gly Ser Ala Pro Asn Phe Pro His Gly 305 310 315 320

Ile Ala Asp Asp Ile Glu Gly Leu Gly Lys Ile Ala Gln Lys Tyr Lys 325 330 335

Leu Pro Leu His Val Asp Ser Cys Leu Gly Ser Phe Ile Val Ser Phe 340 345 350

Met Glu Lys Ala Gly Tyr Lys Asn Leu Pro Leu Leu Asp Phe Arg Val 355 360 365

Pro Gly Val Thr Ser Ile Ser Cys Asp Thr His Lys Tyr Gly Phe Ala 370 375 380

Pro Lys Gly Ser Ser Val Ile Met Tyr Arg Asn Ser Asp Leu Arg Met 385 390 395 400

His Gln Tyr Tyr Val Asn Pro Ala Trp Thr Gly Gly Leu Tyr Gly Ser 405 410 415

Pro Thr Leu Ala Gly Ser Arg Pro Gly Ala Ile Val Val Gly Cys Trp
420 425 430

Ala Thr Met Val Asn Met Gly Glu Asn Gly Tyr Ile Glu Ser Cys Gln
435 440 445

```
Glu Ile Val Gly Ala Ala Met Lys Phe Lys Lys Tyr Ile Gln Glu Asn
    450
                        455
Ile Pro Asp Leu Asn Ile Met Gly Asn Pro Arg Tyr Ser Val Ile Ser
465
                    470
                                         475
                                                             480
Phe Ser Ser Lys Thr Leu Asn Ile His Glu Leu Ser Asp Arg Leu Ser
                485
                                     490
                                                         495
Lys Lys Gly Trp His Phe Asn Ala Leu Gln Lys Pro Val Ala Leu His
            500
                                 505
Met Ala Phe Thr Arg Leu Ser Ala His Val Val Asp Glu Ile Cys Asp
        515
                             520
                                                 525
Ile Leu Arg Thr Thr Val Gln Glu Leu Lys Ser Glu Ser Asn Ser Lys
                        535
                                             540
Pro Ser Pro Asp Gly Thr Ser Ala Leu Tyr Gly Val Ala Gly Ser Val
545
                    550
                                         555
                                                              560
Lys Thr Ala Gly Val Ala Asp Lys Leu Ile Val Gly Phe Leu Asp Ala
                565
                                     570
                                                         575
Leu Tyr Lys Leu Gly Pro Gly Glu Asp Thr Ala Thr Lys
            580
                                 585
```

<211> 3470

<212> DNA

<213> Saccharomyces cerevisiae

<400> 59

acgtcatttt gttcttgtgg agctggtgt tcttgtggag cagattcctg tggagcagat 60 tcctgtggaa cttgtggttc ttgtggagca tctggagcct gtggtggttc ttggtgctgt 120 tcttcgactg gggcatcgac aacagattct ggtagttgct ctacgtgagt ttcttgagct 180 tcagacatta tccttatggt tttagcgtaa ttgcttaatt ttgattcctt caaaagtata 240 tatatttaga agagaggaaa ttattttct catgtccttt ttaaatccct ttgggtggcg 300 aaaaaaaaaga atgtaaaaaa ttttgccctt cgtttacagt gataaatata cggaggggct 360 ctatgataaa ggtagtagta aatcattgaa ttgttgaaca agcattgaca gatatgataa 420 caagcaattg taatcaataa gccaccaatt agaaggctac tcaaaagaat aaagttacta 480 taaaatatac tgcggtatat atgttgcggt ttcagcgatt tgcgtcctcg tatgcccaag 540 cacaggccgt tagaaaatat ccagttggag gtatattcca tggttatgaa gtgagaagaa 600 tcctaccggt tccggagctg agactcactg cggtagattt ggtgcactcc cagacaggag 660 ccgagcattt gcatattgat agagacgaca agaataatgt gttcagcatt gcttttaaaa 720

```
ccaaccetec agatteeact ggggteecte atattetaga gcatacaacg ttgtgtgggt 780
ctgttaaata tccagttagg gacccttttt tcaaaatgct aaataaatct ctagctaatt 840
tcatgaacgc tatgacaggt ccagattata cattttttcc cttttccact acgaaccctc 900
aagatttege taatttaaga ggtgtttatt tagacteeac ettgaateeg etaettaaac 960
aagaagattt tgatcaggag ggttggaggt tggagcataa aaacatcaca gacccggaga 1020
gtaacattgt tttcaaaggt gttgtctata acgaaatgaa aggtcaaata tcaaatgcca 1080
attactattt ctggagtaaa tttcaacagt ctatttatcc ttccctgaat aactccggcg 1140
gagatectat gaaaattaca gaettgagat aeggegatet ettggattte eateacaaaa 1200
attaccatcc ctccaatgca aaaactttca cgtacggtaa cttgccattg gtggatacgt 1260
taaagcaatt aaatgagcag ttcagtggtt acgggaagag agctcgaaag gataagttgt 1320
taatgcctat tgatttaaaa aaagacatag atgtcaagtt actgggtcaa atagatacta 1380
tgcttccacc ggagaagcag acaaaagcct caatgacgtg gatttgtgga gcgccacagg 1440
acacatatga tacctttttg ttaaaagtac tggggaattt attaatggat ggccattctt 1500
ctgtaatgta tcaaaaatta atagaatcag gaattggttt ggagttctcc gtaaattcag 1560
gtgttgaacc aactacagca gtaaatttgc taactgttgg tatacagggc gtgagtgata 1620
ttgaaatatt taaagacact gtaaataata tttttcaaaa cctgttggaa acagaacatc 1680
cttttgaccg caagcgtatc gatgccataa ttgaacaatt ggaattatct aagaaggatc 1740
aaaaggctga ctttggactt caattactct attctatact acctggttgg acaaacaaaa 1800
tcgatccttt tgagagcttg ttgtttgagg acgttttgca aagatttaga ggtgacttag 1860
aaacgaaagg tgatacttta ttccaagatt taatccgtaa atatatcgtt cataaacctt 1920
gtttcacgtt ttccattcag ggatctgaag agttctctaa atctttggat gatgaagaac 1980
aaacaagact gagagaaaaa attactgcct tggatgaaca agacaagaaa aacatcttta 2040
aacgtggtat actgttacag gagaaacaaa atgaaaaaga agatttatcc tgtttaccta 2100
ccttacaaat aaaagacatc ccaagagctg gtgataaata ttcaatcgaa cagaagaata 2160
atacaatgtc taggattact gataccaatg gtatcacata tgtcagaggt aaacgtttac 2220
taaatgacat aataccettt gaactettee catacttace tttatttget gaategttaa 2280
ctaacctagg gacaacaaca gaatccttca gtgaaataga agatcagata aaattacata 2340
egggtggtat atcaacacat gtagaggtta catctgacce taacaccaca gagcetegce 2400
tgattttcgg gtttgacgga tggtctttaa attcgaagac cgaccacatt tttgaattct 2460
ggtctaagat cttactagaa actgatttcc ataaaaacag cgataaattg aaagttctta 2520
tccgcttatt agcatettca aacacatett ctgtagcaga tgccggtcat gcatttgcaa 2580
ggggctattc tgccgcacat tatagatcaa gtggagctat aaatgagacc ctcaatggta 2640
ttgagcaact acaatttata aatagattgc acagcttgtt agacaatgaa gaaactttcc 2700
aaagagaagt tgtcgacaag ctaactgaat tgcaaaagta cattgttgat accaataaca 2760
tgaatttttt tatcacctca gactctgatg ttcaagcgaa aacagtagaa agccaaattt 2820
caaaattcat ggagagatta cctcatggca gctgcttgcc caatggacca aagacttcag 2880
attatectet tattggatee aaatgtaaac ataetttgat aaaattteet ttecaggtee 2940
attacacatc ccaagcttta ttgggtgtgc cgtatacaca taaggatggc tctgcacttc 3000
aagttatgtc aaatatgcta acattcaaac atttgcacag agaagtcaga gaaaagggtg 3060
gtgcttatgg tggtggtgct tcttatagcg ccttagcggg tattttcagt ttctattcct 3120
atagggatcc tcagcctttg aagagtttag aaaccttcaa gaatagcggg cgttatatac 3180
tgaacgatgc caagtggggc gtcacagacc ttgatgaagc taaattgaca atatttcaac 3240
aagtagacgc acctaaaagt cccaaaggag aaggcgtgac gtatttcatg agcggtgtta 3300
cagacgatat gaaacaagca agaagggaac aactcttaga cgtatctctc ctggacgttc 3360
atagagtcgc cgaaaaatat ctactaaaca aagaaggggt gagtacggtc attggacctg 3420
```

qaatcgaggg gaagactgtt tcaccaaatt gggaggtgaa ggaactgtag

<211> 989

<212> PRT

<213> Saccharomyces cerevisiae

<400> 60

Met Leu Arg Phe Gln Arg Phe Ala Ser Ser Tyr Ala Gln Ala Gln Ala 1 5 10 15

Val Arg Lys Tyr Pro Val Gly Gly Ile Phe His Gly Tyr Glu Val Arg
20 25 30

Arg Ile Leu Pro Val Pro Glu Leu Arg Leu Thr Ala Val Asp Leu Val
35 40 45

His Ser Gln Thr Gly Ala Glu His Leu His Ile Asp Arg Asp Asp Lys 50 55 60

Asn Asn Val Phe Ser Ile Ala Phe Lys Thr Asn Pro Pro Asp Ser Thr 65 70 75 80

Gly Val Pro His Ile Leu Glu His Thr Thr Leu Cys Gly Ser Val Lys 85 90 95

Tyr Pro Val Arg Asp Pro Phe Phe Lys Met Leu Asn Lys Ser Leu Ala 100 105 110

Asn Phe Met Asn Ala Met Thr Gly Pro Asp Tyr Thr Phe Phe Pro Phe 115 120 125

Ser Thr Thr Asn Pro Gln Asp Phe Ala Asn Leu Arg Gly Val Tyr Leu 130 135 140

Asp Ser Thr Leu Asn Pro Leu Leu Lys Gln Glu Asp Phe Asp Gln Glu 145 150 155 160

Gly Trp Arg Leu Glu His Lys Asn Ile Thr Asp Pro Glu Ser Asn Ile 165 170 175

Val Phe Lys Gly Val Val Tyr Asn Glu Met Lys Gly Gln Ile Ser Asn 180 185 190

Ala Asn Tyr Tyr Phe Trp Ser Lys Phe Gln Gln Ser Ile Tyr Pro Ser 195 200 205

Leu Asn Asn Ser Gly Gly Asp Pro Met Lys Ile Thr Asp Leu Arg Tyr 210 215 220

Gly Asp Leu Leu Asp Phe His His Lys Asn Tyr His Pro Ser Asn Ala 225 230 230 235 235

Lys Thr Phe Thr Tyr Gly Asn Leu Pro Leu Val Asp Thr Leu Lys Gln 245 250 255

Leu Asn Glu Gln Phe Ser Gly Tyr Gly Lys Arg Ala Arg Lys Asp Lys 260 265 270

Leu Leu Met Pro Ile Asp Leu Lys Lys Asp Ile Asp Val Lys Leu Leu 275 280 285

Gly Gln Ile Asp Thr Met Leu Pro Pro Glu Lys Gln Thr Lys Ala Ser 290 295 300

Met Thr Trp Ile Cys Gly Ala Pro Gln Asp Thr Tyr Asp Thr Phe Leu 305 310 315 320

Leu Lys Val Leu Gly Asn Leu Leu Met Asp Gly His Ser Ser Val Met 325 330 335

Tyr Gln Lys Leu Ile Glu Ser Gly Ile Gly Leu Glu Phe Ser Val Asn 340 345 350

Ser Gly Val Glu Pro Thr Thr Ala Val Asn Leu Leu Thr Val Gly Ile 355 360 365

Gln Gly Val Ser Asp Ile Glu Ile Phe Lys Asp Thr Val Asn Asn Ile 370 375 380

Phe Gln Asn Leu Leu Glu Thr Glu His Pro Phe Asp Arg Lys Arg Ile 385 390 395 400

Asp Ala Ile Ile Glu Gln Leu Glu Leu Ser Lys Lys Asp Gln Lys Ala 405 410 415

Asp Phe Gly Leu Gln Leu Leu Tyr Ser Ile Leu Pro Gly Trp Thr Asn 420 425 430

Lys Ile Asp Pro Phe Glu Ser Leu Leu Phe Glu Asp Val Leu Gln Arg 435 440 445

Phe Arg Gly Asp Leu Glu Thr Lys Gly Asp Thr Leu Phe Gln Asp Leu 450 455 460

Ile Arg Lys Tyr Ile Val His Lys Pro Cys Phe Thr Phe Ser Ile Gln 470 475 480

Gly Ser Glu Glu Phe Ser Lys Ser Leu Asp Asp Glu Glu Gln Thr Arg 485 490 495

Leu Arg Glu Lys Ile Thr Ala Leu Asp Glu Gln Asp Lys Lys Asn Ile 500 505 510

Phe Lys Arg Gly Ile Leu Leu Gln Glu Lys Gln Asn Glu Lys Glu Asp 515 520 525

Leu Ser Cys Leu Pro Thr Leu Gln Ile Lys Asp Ile Pro Arg Ala Gly 530 535 540

Asp Lys Tyr Ser Ile Glu Gln Lys Asn Asn Thr Met Ser Arg Ile Thr 545 550 555 560

Asp Thr Asn Gly Ile Thr Tyr Val Arg Gly Lys Arg Leu Leu Asn Asp 565 570 575

Ile Ile Pro Phe Glu Leu Phe Pro Tyr Leu Pro Leu Phe Ala Glu Ser 580 585 590

Leu Thr Asn Leu Gly Thr Thr Thr Glu Ser Phe Ser Glu Ile Glu Asp 595 600 605

Gln Ile Lys Leu His Thr Gly Gly Ile Ser Thr His Val Glu Val Thr 610 615 620

Ser Asp Pro Asn Thr Thr Glu Pro Arg Leu Ile Phe Gly Phe Asp Gly 625 630 635 640

Trp Ser Leu Asn Ser Lys Thr Asp His Ile Phe Glu Phe Trp Ser Lys 645 650 655

Ile Leu Leu Glu Thr Asp Phe His Lys Asn Ser Asp Lys Leu Lys Val 660 665 670

Leu Ile Arg Leu Leu Ala Ser Ser Asn Thr Ser Ser Val Ala Asp Ala 675 680 685

Gly His Ala Phe Ala Arg Gly Tyr Ser Ala Ala His Tyr Arg Ser Ser 690 695 700

Gly Ala Ile Asn Glu Thr Leu Asn Gly Ile Glu Gln Leu Gln Phe Ile 705 710 715 720

Asn Arg Leu His Ser Leu Leu Asp Asn Glu Glu Thr Phe Gln Arg Glu
725 730 735

Val Val Asp Lys Leu Thr Glu Leu Gln Lys Tyr Ile Val Asp Thr Asn 740 745 750

Asn Met Asn Phe Phe Ile Thr Ser Asp Ser Asp Val Gln Ala Lys Thr 755 760 765

Val Glu Ser Gln Ile Ser Lys Phe Met Glu Arg Leu Pro His Gly Ser 770 775 780

Cys Leu Pro Asn Gly Pro Lys Thr Ser Asp Tyr Pro Leu Ile Gly Ser 785 790 795 800

Lys Cys Lys His Thr Leu Ile Lys Phe Pro Phe Gln Val His Tyr Thr 805 810 815

Ser Gln Ala Leu Leu Gly Val Pro Tyr Thr His Lys Asp Gly Ser Ala 820 825 830

Leu Gln Val Met Ser Asn Met Leu Thr Phe Lys His Leu His Arg Glu 835 840 845

Val Arg Glu Lys Gly Gly Ala Tyr Gly Gly Gly Ala Ser Tyr Ser Ala 850 855 860

Leu Ala Gly Ile Phe Ser Phe Tyr Ser Tyr Arg Asp Pro Gln Pro Leu 865 870 875 880

Lys Ser Leu Glu Thr Phe Lys Asn Ser Gly Arg Tyr Ile Leu Asn Asp 885 890 895

Ala Lys Trp Gly Val Thr Asp Leu Asp Glu Ala Lys Leu Thr Ile Phe 900 905 910

Gln Gln Val Asp Ala Pro Lys Ser Pro Lys Gly Glu Gly Val Thr Tyr 915 920 925

Phe Met Ser Gly Val Thr Asp Asp Met Lys Gln Ala Arg Arg Glu Gln 930 935 940

Leu Leu Asp Val Ser Leu Leu Asp Val His Arg Val Ala Glu Lys Tyr 945 950 955 960

Leu Leu Asn Lys Glu Gly Val Ser Thr Val Ile Gly Pro Gly Ile Glu 965 970 975

Gly Lys Thr Val Ser Pro Asn Trp Glu Val Lys Glu Leu 980 985

```
<210> 61
<211> 1612
<212> DNA
<213> Saccharomyces cerevisiae
<400> 61
cttttctcag cacctgtcca gagacataac atcacaatca catcgcccca gtaaatgcat 60
acgcaagata agatacaaac tggctacggg aacactacgc caacgtgatt ggcaattgtg 120
ctctaatagt tactctatta ttgctgttaa ttgacaatgt ttagtcacgt gcaacacaat 180
tcaagtcacg tggaaaggcc ttcacatggt gatccatctt ctacatcttc atcggtcctg 240
cataaagtca taatatgggg ctactggaat gtatgcactt aacagtacta ttatatggtg 300
aggetgtaat gettacegtt ttgtggetat tetegtatte tgtaggeece eecatacaca 360
tttttcggta actgcggcat atagatgaaa gttgaaatga atattcaaaa gaatatatat 420
aataatgcag gagatcaagg aagaattaga tatgtataag agtgatggta gaggcaaaaa 480
ataaaaagta agcaggagaa tgaatcgtgt tggtatagac gtagatcata tgataggggt 540
cctgcttctg gccgtagtgg tggtgttttg ggttggcgct tcgtgtttga ctaatgaatt 600
gctcgagaca aacgcgtaca ataaaccttt cttccttact tatctaaaca tatcatcgtt 660
tgctctttat ttgacgccag atctatggag gataatccaa tcaagaagga agagcttgca 720
ggaacggaca gaacgaacat tacctattca cacacaagaa tctttttcag agttcctacc 780
tttactatct tcaactcctt ctacttcttc aaatttgtct tcgatagcgg acacgaaagt 840
gaaggataca atgaggttga gtctgctatt ttgcgtcttg tggttcgtgg caaatttggc 900
ggctaacgct gctttgtcgt ataccacagt ggcttcgtca acaattcttt catcgacatc 960
ctcatttttt accttatttc ttgccactag tctaggaata gaaacttttt cgacaaaaaa 1020
actgctgggg ttatttgtgt ctttgtttgg aattatctta attgtgatgc aatcctcgaa 1080
gcaacaggat tctgtgagtg cttcctcctt tttggtaggt aacactttag cactgctggg 1140
gtcattgggt tacagtgtct atacaaccct tttgaaatac gaaatatcat ccaaaggtct 1200
cagactagac attcagatgt ttcttggtta tgttggtatc ttcacgtttc tgttgttttg 1260
gccaatttta ataatcctgg atataacaca tatggaaact tttgaactac caagtaactt 1320
ccacatttct tttcttgtca tgttaaattg tatcattatc tttgttagtg actatttttg 1380
gtgtaaagcc ctcattttga catcaccctt ggtggttacc gttgccttaa cttttactat 1440
cccgttagcc atgttcgctg attttgtatg gcgagaggca ttttttacgc cttggtatat 1500
cattggtgtt attttcattt ttgtttcatt ctttctagtt aaccatcggg gagaatctgc 1560
tgttgaaaag gactgtgctg cggttgaaaa aggacctatc ttggatgcct aa
<210> 62
<211> 370
<212> PRT
<213> Saccharomyces cerevisiae
<400> 62
Met Asn Arg Val Gly Ile Asp Val Asp His Met Ile Gly Val Leu Leu
                                     10
Leu Ala Val Val Val Phe Trp Val Gly Ala Ser Cys Leu Thr Asn
                                 25
             20
```

Glu	Leu	Leu	Glu	Thr	Asn	Ala	Tyr	Asn	Lys	Pro	Phe	Phe	Leu	Thr	Туг
		35					40					45			

Leu Asn Ile Ser Ser Phe Ala Leu Tyr Leu Thr Pro Asp Leu Trp Arg
50 55 60

Ile Ile Gln Ser Arg Arg Lys Ser Leu Gln Glu Arg Thr Glu Arg Thr 65 70 75 80

Leu Pro Ile His Thr Gln Glu Ser Phe Ser Glu Phe Leu Pro Leu Leu 85 90 95

Ser Ser Thr Pro Ser Thr Ser Ser Asn Leu Ser Ser Ile Ala Asp Thr 100 105 110

Lys Val Lys Asp Thr Met Arg Leu Ser Leu Leu Phe Cys Val Leu Trp 115 120 125

Phe Val Ala Asn Leu Ala Ala Asn Ala Ala Leu Ser Tyr Thr Thr Val 130 135 140

Ala Ser Ser Thr Ile Leu Ser Ser Thr Ser Ser Phe Phe Thr Leu Phe 145 150 155 160

Leu Ala Thr Ser Leu Gly Ile Glu Thr Phe Ser Thr Lys Lys Leu Leu 165 170 175

Gly Leu Phe Val Ser Leu Phe Gly Ile Ile Leu Ile Val Met Gln Ser 180 185 190

Ser Lys Gln Gln Asp Ser Val Ser Ala Ser Ser Phe Leu Val Gly Asn 195 200 205

Thr Leu Ala Leu Leu Gly Ser Leu Gly Tyr Ser Val Tyr Thr Thr Leu 210 215 220

Leu Lys Tyr Glu Ile Ser Ser Lys Gly Leu Arg Leu Asp Ile Gln Met 225 230 235 240

Phe Leu Gly Tyr Val Gly Ile Phe Thr Phe Leu Leu Phe Trp Pro Ile 245 250 255

Leu Ile Ile Leu Asp Ile Thr His Met Glu Thr Phe Glu Leu Pro Ser 260 265 270

Asn Phe His Ile Ser Phe Leu Val Met Leu Asn Cys Ile Ile Ile Phe 275 280 285

```
Val Ser Asp Tyr Phe Trp Cys Lys Ala Leu Ile Leu Thr Ser Pro Leu
    290
                        295
                                            300
Val Val Thr Val Ala Leu Thr Phe Thr Ile Pro Leu Ala Met Phe Ala
305
                    310
                                        315
                                                             320
Asp Phe Val Trp Arg Glu Ala Phe Phe Thr Pro Trp Tyr Ile Ile Gly
                325
                                    330
                                                        335
Val Ile Phe Ile Phe Val Ser Phe Phe Leu Val Asn His Arg Gly Glu
            340
                                345
                                                    350
Ser Ala Val Glu Lys Asp Cys Ala Ala Val Glu Lys Gly Pro Ile Leu
                            360
                                                365
Asp Ala
    370
<210> 63
<211> 1376
<212> DNA
<213> Saccharomyces cerevisiae
<400> 63
cggcccatga gcataaattg agaacgatat tattaattct agtatatttt aggaacagca 60
ggccattata tccaaaactt ttttaaaata gcatccacat cacttactgg cgtacagttt 120
cgtgtctggc aacaaaaag tacatttaat ttcaatgatt aaaaagacat taaacatccg 180
tacattacgc acccatgcat gctatctgaa aatattctac atgctgcttt tagaaatttg 240
aagcggcata tggtgtttcc ttgcggagac gcgcgctcag gggaactgcg attccgaaag 300
atgeetttee cagteecata tgeecateee aagateatge eegggagage aaatgtegee 360
ccagccaggt cggacacatc tgtcattcga ccactaagta ttgtcaaagc tagggttaat 420
tgaagatagc tctacatgtt attagtagag tttttaaacg ttgagatact agtgaacgta 480
tacacaagag cggataaaag atgtctttag ttgtccaaga acaaggttcc ttccaacaca 540
ttttacggta tgtttattat tactagtgag ctatgacaaa atcggctaaa aacttaaaaa 600
tatgacagac aagaaaggaa attcattcac ctctttaatc gtggcgatta ttcgctaacg 660
acgtgattaa atataataat gatgttcccg ggctggacta agaacgtaat aataaggctg 720
atgaacaaat ttatggttta tgcatcaaaa aggaacatga atttggcata agcgcacata 780
attacggaac attgcaatag cggtcggagg taaagtaccg ggaatgcttt acaatgaatc 840
```

agctattggc ggatattgac caaccaacac taaaatttta tatcctgctc cgattttgt 900 cttggccgtg aaatccatta tgcacatttt ttactaacgt ttatcaataa gttcggtttc 960 ccgtctaaat tttttacgc agtttgttga acactaacgt tgacggtaac attaagatcg 1020 tttacgcttt gaccactatc aagggtgttg gtcgtcgtta ctccaacttg gtctgtaaga 1080 aggctgatgt tgattacac aagagagctg gtgaattgac ccaagaagaa ttggaaagaa 1140 ttgttcaaat tatgcaaaac ccaactcact acaaaatccc agcttggttc ttgaaccgtc 1200 aaaatgacat cactgatggt aaggactacc acactttggc taacaacgtc gaatccaaat 1260



tgagagatga cttggaaaga ttaaagaaaa tcagagctca tcgtggtatc agacacttct 1320 ggggtttgcg tgttagaggt caacacacca agaccactgg tagaagaaga gcttaa 1376

<210> 64

<211> 146

<212> PRT

<213> Saccharomyces cerevisiae

<400> 64

Met Ser Leu Val Val Gln Glu Gln Gly Ser Phe Gln His Ile Leu Arg

1 5 10 15

Leu Leu Asn Thr Asn Val Asp Gly Asn Ile Lys Ile Val Tyr Ala Leu 20 25 30

Thr Thr Ile Lys Gly Val Gly Arg Arg Tyr Ser Asn Leu Val Cys Lys
35 40 45

Lys Ala Asp Val Asp Leu His Lys Arg Ala Gly Glu Leu Thr Gln Glu 50 55 60

Glu Leu Glu Arg Ile Val Gln Ile Met Gln Asn Pro Thr His Tyr Lys
65 70 75 80

Ile Pro Ala Trp Phe Leu Asn Arg Gln Asn Asp Ile Thr Asp Gly Lys

85 90 95

Asp Tyr His Thr Leu Ala Asn Asn Val Glu Ser Lys Leu Arg Asp Asp 100 105 110

Leu Glu Arg Leu Lys Lys Ile Arg Ala His Arg Gly Ile Arg His Phe 115 120 125

Trp Gly Leu Arg Val Arg Gly Gln His Thr Lys Thr Thr Gly Arg Arg 130 135 140

Arg Ala

145

<210> 65

<211> 1289

<212> DNA

<213> Saccharomyces cerevisiae

<400> 65

```
actgcataca caataactgt agatgtagcc caaggcacta ccacaggtat ttctgctcac 60
gacaggtcga tgacttgtag ggctcttgca gactcttcct ctacgccaaa atcattttta 120
aaaccagggc acatctgtcc cttgagagcc gctgatggcg gtgttttgca gagaagaggc 180
cacactgagg ccggtgtcga tttgtgtaaa ctaagtggac taagtcccgt cgctgttatt 240
ggcgaattgg ttaacgatga cgaacaagga actatgatga gattaaatga ctgccaagcg 300
tttggtaaga aacatggcat tcctttgatc tccatcgaag aattggccca atatttgaag 360
aaataatctg gtgaacattt tctccattca ttctatcaca acagactcac acatatatac 420
atgtatatat ttgtaacttt gtatatatct tttgtttttt gacctttttc ttcctctatg 480
tttttcagcc atacaaaaat atgggatttt tagcaagaga aaaagtacat ctaaaaaaaag 540
tagtaatagg aggaagccaa gattggttga aacacagtta taaactcttc aaggcaatta 600
tgaacaggat tttcggatat gggaacaaaa agagccatga tcagctctta caagagtcga 660
atcagtccat gaatcaggcc caacaatcac tatcgaacag aatatcccag ttagatactc 720
aaatcgccca gttaaacttc cagctgcaaa atattcaaaa gaatttgcaa agatcaaaca 780
acaagcaacc ctcgttaaga aaacaggctt tgaagatttt aaataaacgt aaacagttag 840
aaaatatgaa ggattetta gatteteaat eetggteeat gaegeaagee eagttaacaa 900
atgataactt acagaacaca atgatcacta taaacgcact aaagcaaaca aacaatgcca 960
tgaaggctca atacggcaag ataaatatcg acaaactaca ggacatgcag gatgagatgc 1020
tggatttaat agaacaaggg gatgagctgc aagaagtctt ggcaatgaat aataacagtg 1080
gegagetega egacattagt gatgeagage tggatgeaga getggatget etggeacaag 1140
aggatttcac tttgccaacc agcgaaaact cattaggtaa cgatatgccc agttacttac 1200
taggtgcgaa tgcgccaccg gcttttattg atgaagagcc aaacttagat actgaagaca 1260
aaaataaagc tttagaaagc gctcagtga
                                                                  1289
```

<211> 262

<212> PRT

<213> Saccharomyces cerevisiae

<400> 66

Met Gly Phe Leu Ala Arg Glu Lys Val His Leu Lys Lys Val Val Ile 1 5 10 15

Gly Gly Ser Gln Asp Trp Leu Lys His Ser Tyr Lys Leu Phe Lys Ala

Ile Met Asn Arg Ile Phe Gly Tyr Gly Asn Lys Lys Ser His Asp Gln
35 40 45

Leu Leu Gln Glu Ser Asn Gln Ser Met Asn Gln Ala Gln Gln Ser Leu 50 55 60

Ser Asn Arg Ile Ser Gln Leu Asp Thr Gln Ile Ala Gln Leu Asn Phe 65 70 75 80

Gln Leu Gln Asn Ile Gln Lys Asn Leu Gln Arg Ser Asn Asn Lys Gln 85 90 95

## EP99870141.1



Pro Ser Leu Arg Lys Gln Ala Leu Lys Ile Leu Asn Lys Arg Lys Gln 100 105 110

Leu Glu Asn Met Lys Asp Ser Leu Asp Ser Gln Ser Trp Ser Met Thr 115 120 125

Gln Ala Gln Leu Thr Asn Asp Asn Leu Gln Asn Thr Met Ile Thr Ile 130 135 140

Asn Ala Leu Lys Gln Thr Asn Asn Ala Met Lys Ala Gln Tyr Gly Lys 145 150 · 155 160

Ile Asn Ile Asp Lys Leu Gln Asp Met Gln Asp Glu Met Leu Asp Leu 165 170 175

Ile Glu Gln Gly Asp Glu Leu Gln Glu Val Leu Ala Met Asn Asn Asn 180 185 190

Ser Gly Glu Leu Asp Asp Ile Ser Asp Ala Glu Leu Asp Ala Glu Leu 195 200 205

Asp Ala Leu Ala Gln Glu Asp Phe Thr Leu Pro Thr Ser Glu Asn Ser 210 215 220

Leu Gly Asn Asp Met Pro Ser Tyr Leu Leu Gly Ala Asn Ala Pro Pro 225 230 235 240

Ala Phe Ile Asp Glu Glu Pro Asn Leu Asp Thr Glu Asp Lys Asn Lys 245 250 255

Ala Leu Glu Ser Ala Gln 260

<210> 67

<211> 1295

<212> DNA

<213> Saccharomyces cerevisiae

<400> 67

ggcaaatcaa ttaaaatcc ttttctctc acctttgcta atattaaaaa ccatagttgt 60 aaagggtact taatgctata ttcctgttaa gtttcctatt tcaccttgtt ttttccaatt 120 cttaccaatt tgaagactat gtttttaaac acccaatcat tttccaccca cacatatatt 180 accctttttt tgggtgaaga gaagtagtat tttgttttc atgggagtgg aagtcctttc 240 aaaactaatc cgagcagtag tgctgtctag gcggagatta ttgaaagtcg gcattggctg 300 cagctagcgt tttgtttttg gtactacctg tcaaacccgg cgtctgccta gattgcgcgg 360 agggtacgtt gaaacttttg cctttccacg tcagtttata atatcaaaag cagcaatata 420

-85

```
cacatttaat ggtttgccct tagtgactat tagggcgttt tggtgaaaag aaagagtcgc 480
tcaaagaaat caatataacc atggctaaat ttttgaaagc aggtaaagtt ggtacgtaaa 540
tttaacagag caaacgctct aattaagata tcgaataaaa ggggactttc aggtgcataa 600
gatgggaaat tgtacaatct gagggacaaa caatatggaa agcagtacat gattgtattg 660
tcattgtagg aggtaccaat tgtgccagta aaaaagagag ggcccgggga gcaccacatt 720
cacgaccaat cgtgtcgatc tgcaaaaaag gcacatggaa aatgtaattt atcagtgttc 780
aacactgagt tgaacagata gctatctaat caatgttaac tttccaggga acaaaaagct 840
aaatccactt ctcttttatt tcaaaatatc attagaaata gaaaaattta ctaacaaatt 900
ttcgtattat cgtagctgtc gttgttcgtg gtcgttacgc tggtaagaag gttgtgatcg 960
ttaagccaca cgatgagggt tctaaatctc atccattcgg tcacgctttg gttgccggta 1020
ttgaaaggta cccatcaaag gtcaccaaga agcacggtgc caagaaggtc gctaagagaa 1080
ctaaaatcaa gccattcatc aaagtcgtca actacaacca cttattgcca accagataca 1140
cettggatgt tgaagettte aagagegttg tatetacaga gaettttgaa caaccatece 1200
aacgtgaaga agccaagaag gttgtaaaga aggcatttga agaaagacat caagctggta 1260
aaaaccaatg gttcttctcc aagttgagat tctaa
<210> 68
```

<211> 136

<212> PRT

<213> Saccharomyces cerevisiae

<400> 68

Met Ala Lys Phe Leu Lys Ala Gly Lys Val Ala Val Val Arg Gly
1 5 10 15

Arg Tyr Ala Gly Lys Lys Val Val Ile Val Lys Pro His Asp Glu Gly 20 25 30

Ser Lys Ser His Pro Phe Gly His Ala Leu Val Ala Gly Ile Glu Arg 35 40 45

Tyr Pro Ser Lys Val Thr Lys Lys His Gly Ala Lys Lys Val Ala Lys
50 55 60

Arg Thr Lys Ile Lys Pro Phe Ile Lys Val Val Asn Tyr Asn His Leu 65 70 75 80

Leu Pro Thr Arg Tyr Thr Leu Asp Val Glu Ala Phe Lys Ser Val Val 85 90 95

Ser Thr Glu Thr Phe Glu Gln Pro Ser Gln Arg Glu Glu Ala Lys Lys
100 105 110

Val Val Lys Lys Ala Phe Glu Glu Arg His Gln Ala Gly Lys Asn Gln 115 120 125

Trp Phe Phe Ser Lys Leu Arg Phe

EP99870141.



130 135

```
<210> 69
<211> 2744
<212> DNA
<213> Saccharomyces cerevisiae
```

## <400> 69

```
taacaataga aaaaaataca cacacattag attggaatta gagcttaagt ggtacaaact 60
agggctaata aagaggtaac ggtcggttct ctactaaggt tcgtattgtg tggcaccgat 120
gttaagcact tttaagcgga ataactcgag tggaaatttt atgtttagtt aggttttacc 180
ttgaattttt ttaaaaaaaa aagagtcaga caggctcgct ctttcctact aaatattagg 240
agcaaagcag taaaaagtct ctgaataagg atagtaacct gtagtaactc caaaatttat 300
cttacaaaga gctattagta tcttggttct tctattttct tcgattattg gtgatttttc 360
ccgcctctag ccaaatccga gcgttccatc gattttttgg ggaaaacagc acatgcaata 420
aaataaaaag caaacaaata cgcgatagtg cacgaaacgt caacacaatc atcaaactct 480
tttgcatatt tctattatag atgagacgag aaacggtggg tgaattttct tcagatgacg 540
atgatgatat tetteteggaa ttaggeacca ggeeteeaag gtttaeteaa atacegeeat 600
catcagcagc attacaaaca caaattccca ctactttgga ggttacaacg accacattaa 660
acaataaaca gagtaaaaat gataaccaac tggttaacca actgaataaa gctcaaggtg 720
aagcaagcat gcttcgtgat aaaataaact ttttgaacat tgaaagggaa aaggaaaaga 780
atattcaagc cgtcaaagtt aatgaattgc aagtcaagca tcttcaagag ttggctaaat 840
taaaacaaga attacagaaa ctggaagatg agaagaagtt cttacagatg gaagcgagag 900
gaaaatcgaa aagggaagtt attacgaatg taaaaccacc gtcaacaaca ttatcaacaa 960
acacaaacac tataacgcca gattcgtcct cagttgcaat cgaagcaaaa cctcaatcac 1020
cacaatcaaa aaaacgtaag ataagtgata atttactgaa aaaaaatatg gttcccctaa 1080
acccaaatag gattattccc gatgaaacga gtttatttct agagtcaata ttacttcatc 1140
aaataatagg cgctgacctg agcacaatag aaatattaaa tagattgaag cttgactaca 1200
tcacagagtt taaatttaag aatttcgtca ttgctaaagg agcccccata gggaagtcca 1260
tagtttctct acttttgcga tgtaaaaaga cgttgaccct cgacaggttc atagatactt 1320
tgctagagga tatagctgtt ttgatcaagg aaatatcagt tcatccaaat gaatcgaaat 1380
tagetgtece atttetegtt geattgatgt ateagattgt acaatttegt ectagtgeca 1440
ctcataattt agcactgaag gattgttttc tttttatttg cgatttaata agaatctatc 1500
atcatgtatt gaaagtaccg atacatgaat caaatatgaa tttgcatgta gaacctcaga 1560
ttttccaata tgaactgata gactatttga taatttcgta ttcctttgat ctcctagaag 1620
gtatattaag ggtactgcag tcgcatccta agcaaactta tatggaattt tttgatgaaa 1680
atattctaaa atcatttgaa tttgtctaca aactagcact aaccatttca tacaagccaa 1740
tggtaaatgt aatatttagt gcagtcgagg tcgttaatat tatcactagt ataatattaa 1800
atatggacaa ttcctcagat ctgaaatcct tgataagcgg tagttggtgg agagattgta 1860
ttacaagatt atacgctctt ttggaaaagg aaattaagag tggcgacgta tataatgaaa 1920
atgtggatac tacaactctt catatgtcga agtaccatga cttttttggg ttaatccgaa 1980
atataggtga taatgaattg ggaggattga tatcaaagct gatttatact gaccgattgc 2040
aaagtgtccc aagggtaatt tctaaagagg atattgggat ggatagcgac aaattcactg 2100
cccctataat aggttacaag atggaaaaat ggcttttgaa gttaaaagat gaagttttaa 2160
atatttttga aaatttatta atgatctatg gagacgatgc gaccatagta aatggagaaa 2220
tgctcatcca ctcctctaaa ttcttatcca gggagcaagc gttaatgata gaaaggtacg 2280
```

```
tgggacaaga ctccccgaac ttggacctca gatgccatct tattgaacat actttaacca 2340 taatatatag gctatggaaa gaccatttca aacaattgcg tgaagaacaa atcaagcagg 2400 tagaaagcca attgattatg tcactatgga ggtttctcgt atgccaaacc gaaactgtga 2460 cagcaaacga aagagaaatg agagatcatc gacaccttgt agatagtttg catgatctga 2520 cgataaagga tcaagcctct tattacgaag atgcctttga ggatttacca gaatatatcg 2580 aagaagaatt gaagatgcaa ttgaataaaa gaacggggag aataatgcaa gtaaagtacg 2640 atgagaaatt tcaagaaatg gcaagaacta ttcttgagtc aaaatcattc gatctaacca 2700 cactagagga ggccgattca ttatatatct caatgggact gtaa 2744
```

<211> 747

<212> PRT

<213> Saccharomyces cerevisiae

<400> 70

Met Arg Arg Glu Thr Val Gly Glu Phe Ser Ser Asp Asp Asp Asp Asp 1 10 15

Ile Leu Leu Glu Leu Gly Thr Arg Pro Pro Arg Phe Thr Gln Ile Pro
20 25 30

Pro Ser Ser Ala Ala Leu Gln Thr Gln Ile Pro Thr Thr Leu Glu Val
35 40 45

Thr Thr Thr Leu Asn Asn Lys Gln Ser Lys Asn Asp Asn Gln Leu 50 55 60

Val Asn Gln Leu Asn Lys Ala Gln Gly Glu Ala Ser Met Leu Arg Asp 65 70 75 80

Lys Ile Asn Phe Leu Asn Ile Glu Arg Glu Lys Glu Lys Asn Ile Gln 85 90 95

Ala Val Lys Val Asn Glu Leu Gln Val Lys His Leu Gln Glu Leu Ala 100 105 110

Lys Leu Lys Gln Glu Leu Gln Lys Leu Glu Asp Glu Lys Lys Phe Leu 115 120 125

Gln Met Glu Ala Arg Gly Lys Ser Lys Arg Glu Val Ile Thr Asn Val 130 135 140

Lys Pro Pro Ser Thr Thr Leu Ser Thr Asn Thr Asn Thr Ile Thr Pro 145 150 155 160

Asp Ser Ser Ser Val Ala Ile Glu Ala Lys Pro Gln Ser Pro Gln Ser 165 170 175

Lys Lys Arg Lys Ile Ser Asp Asn Leu Leu Lys Lys Asn Met Val Pro 180 185 190

Leu Asn Pro Asn Arg Ile Ile Pro Asp Glu Thr Ser Leu Phe Leu Glu
195 200 205

Ser Ile Leu Leu His Gln Ile Ile Gly Ala Asp Leu Ser Thr Ile Glu 210 215 220

Ile Leu Asn Arg Leu Lys Leu Asp Tyr Ile Thr Glu Phe Lys Phe Lys 225 230 235 240

Asn Phe Val Ile Ala Lys Gly Ala Pro Ile Gly Lys Ser Ile Val Ser 245 250 255

Leu Leu Leu Arg Cys Lys Lys Thr Leu Thr Leu Asp Arg Phe Ile Asp 260 265 270

Thr Leu Glu Asp Ile Ala Val Leu Ile Lys Glu Ile Ser Val His 275 280 285

Pro Asn Glu Ser Lys Leu Ala Val Pro Phe Leu Val Ala Leu Met Tyr 290 295 300

Gln Ile Val Gln Phe Arg Pro Ser Ala Thr His Asn Leu Ala Leu Lys 305 310 315 320

Asp Cys Phe Leu Phe Ile Cys Asp Leu Ile Arg Ile Tyr His His Val 325 330 335

Leu Lys Val Pro Ile His Glu Ser Asn Met Asn Leu His Val Glu Pro 340 345 350

Gln Ile Phe Gln Tyr Glu Leu Ile Asp Tyr Leu Ile Ile Ser Tyr Ser 355 360 365

Phe Asp Leu Leu Glu Gly Ile Leu Arg Val Leu Gln Ser His Pro Lys 370 375 380

Gln Thr Tyr Met Glu Phe Phe Asp Glu Asn Ile Leu Lys Ser Phe Glu 385 390 395 400

Phe Val Tyr Lys Leu Ala Leu Thr Ile Ser Tyr Lys Pro Met Val Asn 405 410 415

Val Ile Phe Ser Ala Val Glu Val Val Asn Ile Ile Thr Ser Ile Ile 420 425 430

Le	ı Asn	Met	Asp	Asn	Ser	Ser	Asp	Leu	Lys	Ser	Leu	Ile	Ser	Gly	Ser
		435					440					445			

- Trp Trp Arg Asp Cys Ile Thr Arg Leu Tyr Ala Leu Leu Glu Lys Glu 450 455 460
- Ile Lys Ser Gly Asp Val Tyr Asn Glu Asn Val Asp Thr Thr Thr Leu 465 470 475 480
- His Met Ser Lys Tyr His Asp Phe Phe Gly Leu Ile Arg Asn Ile Gly
  485 490 495
- Asp Asn Glu Leu Gly Gly Leu Ile Ser Lys Leu Ile Tyr Thr Asp Arg
  500 505 510
- Leu Gln Ser Val Pro Arg Val Ile Ser Lys Glu Asp Ile Gly Met Asp 515 520 525
- Ser Asp Lys Phe Thr Ala Pro Ile Ile Gly Tyr Lys Met Glu Lys Trp 530 535 540
- Leu Leu Lys Leu Lys Asp Glu Val Leu Asn Ile Phe Glu Asn Leu Leu 545 550 555 560
- Met Ile Tyr Gly Asp Asp Ala Thr Ile Val Asn Gly Glu Met Leu Ile 565 570 575
- His Ser Ser Lys Phe Leu Ser Arg Glu Gln Ala Leu Met Ile Glu Arg 580 585 590
- Tyr Val Gly Gln Asp Ser Pro Asn Leu Asp Leu Arg Cys His Leu Ile 595 600 605
- Glu His Thr Leu Thr Ile Ile Tyr Arg Leu Trp Lys Asp His Phe Lys 610 615 620
- Gln Leu Arg Glu Glu Gln Ile Lys Gln Val Glu Ser Gln Leu Ile Met 625 630 635 640
- Ser Leu Trp Arg Phe Leu Val Cys Gln Thr Glu Thr Val Thr Ala Asn 645 650 655
- Glu Arg Glu Met Arg Asp His Arg His Leu Val Asp Ser Leu His Asp 660 665 670
- Leu Thr Ile Lys Asp Gln Ala Ser Tyr Tyr Glu Asp Ala Phe Glu Asp 675 680 685

```
      Leu Pro Glu Tyr Glu Glu Glu Glu Glu Leu Lys Met Gln Leu Asn Lys Arg 690
      The Gly Arg Ile Met Gln Val Lys Tyr Asp Glu Lys Phe Gln Glu Met 715
      Glu Met 725

      Ala Arg Thr Ile Leu Glu 735
      Ser Lys Ser Phe Asp Leu Thr Thr Leu Glu 735
```

Glu Ala Asp Ser Leu Tyr Ile Ser Met Gly Leu 740 745

<210> 71 <211> 3929 <212> DNA <213> Saccharomyces cerevisiae

<400> 71 ctctttaaat attctatgta ctttgtgcaa acatcattgt catcacataa atgcattcct 60 actattacta acttgaactt cacttcactg gaagaactgg gttattcaag gtaaagaaat 120 catttgtttt gcgccaagtt tcgtctgaca agaatttttt attattattt cccacttttc 180 atcgaaggaa acgcgtcaaa tccattcgtt actacgcgca atctgcgtta tttccttttt 240 cggcatacca tcgcgaaata tcaacggcca caccatagat tccttttgat gttaaattaa 300 aagcgactga atgaagtgca cacattttt atttcttctt gattttcttt tctattttgt 360 tttgctttct cttctgtcga caacgtctca actgtactca ccattagtat tctcgaaggc 420 tttagcagac ttgtgaataa ttaattgccc actttgatca agaaagatat tcgcagcaca 480 atacaataat aacattcaaa atggcaatca atggtaacag tattcctgcc ataaaggata 540 ataccatcgg tccatggaaa ctaggtgaaa ctctcggtct agggagcact ggtaaagtcc 600 agcttgctcg taatggatcc acaggacaag aggcggcagt taaggtaata tcaaaagcag 660 tattcaatac cggtaatgtc agcggtactt cgattgttgg ctccaccacc ccagatgctc 720 taccatatgg tatagaacgc gaaataatca ttatgaagtt gttaaaccac ccaaatgtgt 780 tacgtttata tgatgtctgg gaaacaaata cagatttata ccttgtttta gaatacgcgg 840 agaaaggtga gttgttcaac ttattggttg agagaggtcc tctgccagag catgaagcta 900 tcaggttttt tagacaaatt attattggtg tgtcgtactg tcatgcgttg ggtattgtcc 960 atcgtgatct aaaaccggaa aatctattat tagatcataa atataacatc aagattgcag 1020 attttggtat ggctgctttg gaaactgaag gaaagctact ggagacgtcg tgcggatcac 1080 cacattatgc tgcaccagaa attgtatctg gtataccgta tcaaggtttc gcaagtgatg 1140 tgtggtcatg cggtgtgatc ctattcgccc ttcttactgg tcggttaccc tttgacgagg 1200 aagatggaaa tataagaaca ctattactta aagttcaaaa aggtgagttc gaaatgcctt 1260 ctgatgatga aatttcgcgt gaagctcagg atttgattag aaaaatctta accgttgatc 1320 ctgaaagaag aatcaagacc agagatatac tcaaacatcc gctattacaa aaatatccaa 1380 gtataagaga ttctaaaagt attagaggct taccaagaga agacacatat ctcacgccat 1440 tatcagaaag taattettet attgacgeta egattttgca aaatttagta atattatgge 1500 atggaagaga teetgaagga attaaggaaa aactaagaga acetggeget aatgeagaaa 1560 · agacattata tgcactactg tatagattta agtgtgacac tcaaaaagag cttattaagc 1620

```
aacagcaagt taagaagagg cagtcaatta gtagcgtttc tgtttcceea tctaaaaaag 1680
tatcgacaac tccacaacgc agaagaaata gagaatcttt aattagtgta acatcttctc 1740
gtaaaaagcc aatateette aacaaattea etgeeteeag tgeeteetee agcaatetaa 1800
ctacacccgg ttcttcaaaa cgcctttcaa aaaacttctc ttcaaagaag aaattatcta 1860
caatcgttaa ccaatcttct ccaacaccag catcacgtaa taaaagagct tcggttataa 1920
atgtggaaaa gaatcaaaaa agagcctcta tcttttctac taccaagaag aacaaaagat 1980
cttctagatc tatcaagaga atgtcattga taccaagcat gaaacgtgaa tcggtgacaa 2040
caaaattaat gtcaacatat gcgaaattgg cagaggatga cgattgggaa tacattgaga 2100
aggaaacaaa gagaacgagc tcaaattttg caactttgat agatgaaatt tttgagtacg 2160
aaaagtacga acaaataagg aaagagaagg aagagctaga acgtaaagtg agagaagcaa 2220
aagcacgtga agagctggaa cgtagaagac gtaaacaaga agaaaaagaa cgtgcaagaa 2280
aattactaga aaaggaagat ctgaaaagaa aacaggagga actcaagaag caaattgaaa 2340
ttgatataag tgatctagag caagagctgt ccaaacacaa agaggaaaaa ctggatggta 2400
atattagatc tatctctgct cctatggaaa atgaagagaa aaatatcaat catttggagg 2460
ttgatattga caatattctc cgtcgccgca acttttcttt acaaactaga cctgtgtcaa 2520
ggcttgatcc gggtataatg ttctccagtc caactgagga agtaagtcca gtggaaccaa 2580
agagaacaga aaatgaaaga cttacaacag aaaagaaaat tttagaaact atcagaagat 2640
caaaattctt gggttcatca tttaatatcg ataaagagtt gaaattgtct aaaatggaat 2700
atccaagtat aattgcacca caaagattgt cagaggagcg agtggtgtca gattctaatg 2760
atggatatga atctttgatc ctcccgaagg atgggaatgg cgtatctcaa ttaaaggata 2820
gtaccgcaac aactgctccc gtctctgatg gtaggttgag gaagatctct gaaattagag 2880
taccacaatt tactagaaaa tcaaggcatt ttagtgagtc caataaaagg ctatctgtcc 2940
tgtcgatgta ctctaccaag gagtcgttta ccaacttggt tgatattttg aaaaacggta 3000
accttgatgt caataaccaa caaagccaaa gaattccaac accaagaagt gcggatgatt 3060
cagaatttet ttttgaaact gtcaacgaag aagetgaata tacaggaaat agttcgaacg 3120
atgagagatt gtacgatgtc ggtgattcca ctatcaaaga caaatccgcg ttaaagctga 3180
actttgcaga tcgttttaat ggatcgaacg aagcgaaaca aactgataac ttacatcttc 3240
cgatccttcc tccgcttaat ggtgacaatg aattgcgtaa acagaatagc caagagggtg 3300
accaggcaca tccaaagatt aaatcgatga taccagaatc aggctcttct tcacatactg 3360
aaaaggaaga agaaaatgag gaaaaggaag agaaaaagcc agaacaacac aaacaagaag 3420
aggatcaaga aaaaagagag aaagtagtag atgatatgga gccaccattg aacaaatctg 3480
tgcaaaaaat tagggaaaaa aatgctggct cgcaggcaaa ggatcattca aaagatcact 3540
taaaagagca taagcaggat aaaaatacag caattggaaa tggttccttc tttagaaaat 3600
teteaaaate tteggacaaa acaatggaat tgtatgecaa gatttetgea aaacaattgt 3660
ttaatggttt agagaagctg ttgcgtggtt ggactcagta tggtttaaaa aatataaaat 3720
cgcaccccaa caatctgacc ttaacgggta aactatcgag tgataatata ttctcactac 3780
gttcaacact ctttgaggtt aatatttatc cgagaggtaa gatgagcgtt gtgcagttca 3840
agaaagtttc tggttcattc aaagctgtca aaaagttggt caatgaagtt gagaacgtcc 3900
tgaataagga aggcgttcta caaaaatag
```

```
<210> 72
```

<sup>&</sup>lt;211> 1142

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Saccharomyces cerevisiae

<sup>&</sup>lt;400> 72

Met Ala Ile Asn Gly Asn Ser Ile Pro Ala Ile Lys Asp Asn Thr Ile

Gly Pro Trp Lys Leu Gly Glu Thr Leu Gly Leu Gly Ser Thr Gly Lys

Val Gln Leu Ala Arg Asn Gly Ser Thr Gly Gln Glu Ala Ala Val Lys 

Val Ile Ser Lys Ala Val Phe Asn Thr Gly Asn Val Ser Gly Thr Ser 

Ile Val Gly Ser Thr Thr Pro Asp Ala Leu Pro Tyr Gly Ile Glu Arg 

Glu Ile Ile Met Lys Leu Leu Asn His Pro Asn Val Leu Arg Leu

Tyr Asp Val Trp Glu Thr Asn Thr Asp Leu Tyr Leu Val Leu Glu Tyr 

Ala Glu Lys Gly Glu Leu Phe Asn Leu Leu Val Glu Arg Gly Pro Leu 

Pro Glu His Glu Ala Ile Arg Phe Phe Arg Gln Ile Ile Gly Val 

Ser Tyr Cys His Ala Leu Gly Ile Val His Arg Asp Leu Lys Pro Glu 

Asn Leu Leu Leu Asp His Lys Tyr Asn Ile Lys Ile Ala Asp Phe Gly 

Met Ala Ala Leu Glu Thr Glu Gly Lys Leu Leu Glu Thr Ser Cys Gly 

Ser Pro His Tyr Ala Ala Pro Glu Ile Val Ser Gly Ile Pro Tyr Gln 

Gly Phe Ala Ser Asp Val Trp Ser Cys Gly Val Ile Leu Phe Ala Leu 

Leu Thr Gly Arg Leu Pro Phe Asp Glu Glu Asp Gly Asn Ile Arg Thr 

Leu Leu Lys Val Gln Lys Gly Glu Phe Glu Met Pro Ser Asp Asp 

Glu Ile Ser Arg Glu Ala Gln Asp Leu Ile Arg Lys Ile Leu Thr Val

260 265 27

Asp Pro Glu Arg Arg Ile Lys Thr Arg Asp Ile Leu Lys His Pro Leu 275 280 285

Leu Gln Lys Tyr Pro Ser Ile Arg Asp Ser Lys Ser Ile Arg Gly Leu 290 295 300

Pro Arg Glu Asp Thr Tyr Leu Thr Pro Leu Ser Glu Ser Asn Ser Ser 305 310 315 320

Ile Asp Ala Thr Ile Leu Gln Asn Leu Val Ile Leu Trp His Gly Arg
325 330 335

Asp Pro Glu Gly Ile Lys Glu Lys Leu Arg Glu Pro Gly Ala Asn Ala 340 345 350

Glu Lys Thr Leu Tyr Ala Leu Leu Tyr Arg Phe Lys Cys Asp Thr Gln 355 360 365

Lys Glu Leu Ile Lys Gln Gln Gln Val Lys Lys Arg Gln Ser Ile Ser 370 375 380

Ser Val Ser Val Ser Pro Ser Lys Lys Val Ser Thr Thr Pro Gln Arg 385 390 395 400

Arg Arg Asn Arg Glu Ser Leu Ile Ser Val Thr Ser Ser Arg Lys Lys
405 410 415

Pro Ile Ser Phe Asn Lys Phe Thr Ala Ser Ser Ala Ser Ser Asn 420 425 430

Leu Thr Thr Pro Gly Ser Ser Lys Arg Leu Ser Lys Asn Phe Ser Ser 435 440 445

Lys Lys Leu Ser Thr Ile Val Asn Gln Ser Ser Pro Thr Pro Ala 450 455 460

Ser Arg Asn Lys Arg Ala Ser Val Ile Asn Val Glu Lys Asn Gln Lys 465 470 475 480

Arg Ala Ser Ile Phe Ser Thr Thr Lys Lys Asn Lys Arg Ser Ser Arg
485 490 495

Ser Ile Lys Arg Met Ser Leu Ile Pro Ser Met Lys Arg Glu Ser Val 500 505 510

Thr Thr Lys Leu Met Ser Thr Tyr Ala Lys Leu Ala Glu Asp Asp Asp

520

525

Trp Glu Tyr Ile Glu Lys Glu Thr Lys Arg Thr Ser Ser Asn Phe Ala 530 535 540

Thr Leu Ile Asp Glu Ile Phe Glu Tyr Glu Lys Tyr Glu Gln Ile Arg 545 550 555 560

Lys Glu Lys Glu Glu Leu Glu Arg Lys Val Arg Glu Ala Lys Ala Arg 565 570 . 575

Glu Glu Leu Glu Arg Arg Arg Lys Gln Glu Glu Lys Glu Arg Ala 580 585 590

Arg Lys Leu Leu Glu Lys Glu Asp Leu Lys Arg Lys Gln Glu Glu Leu 595 600 605

Lys Lys Gln Ile Glu Ile Asp Ile Ser Asp Leu Glu Gln Glu Leu Ser 610 620

Lys His Lys Glu Glu Lys Leu Asp Gly Asn Ile Arg Ser Ile Ser Ala 625 630 635 640

Pro Met Glu Asn Glu Glu Lys Asn Ile Asn His Leu Glu Val Asp Ile 645 650 655

Asp Asn Ile Leu Arg Arg Arg Asn Phe Ser Leu Gln Thr Arg Pro Val 660 665 670

Ser Arg Leu Asp Pro Gly Ile Met Phe Ser Ser Pro Thr Glu Glu Val 675 680 685

Ser Pro Val Glu Pro Lys Arg Thr Glu Asn Glu Arg Leu Thr Thr Glu 690 695 700

Lys Lys Ile Leu Glu Thr Ile Arg Arg Ser Lys Phe Leu Gly Ser Ser 705 710 715 720

Phe Asn Ile Asp Lys Glu Leu Lys Leu Ser Lys Met Glu Tyr Pro Ser 725 730 735

Ile Ile Ala Pro Gln Arg Leu Ser Glu Glu Arg Val Val Ser Asp Ser 740 745 750

Asn Asp Gly Tyr Glu Ser Leu Ile Leu Pro Lys Asp Gly Asn Gly Val 755 760 765

Ser Gln Leu Lys Asp Ser Thr Ala Thr Thr Ala Pro Val Ser Asp Gly

770 775 780

Arg Leu Arg Lys Ile Ser Glu Ile Arg Val Pro Gln Phe Thr Arg Lys 785 790 795 800

Ser Arg His Phe Ser Glu Ser Asn Lys Arg Leu Ser Val Leu Ser Met 805 810 815

Tyr Ser Thr Lys Glu Ser Phe Thr Asn Leu Val Asp Ile Leu Lys Asn 820 825 830

Gly Asn Leu Asp Val Asn Asn Gln Gln Ser Gln Arg Ile Pro Thr Pro 835 840 845

Arg Ser Ala Asp Asp Ser Glu Phe Leu Phe Glu Thr Val Asn Glu Glu 850 855 860

Ala Glu Tyr Thr Gly Asn Ser Ser Asn Asp Glu Arg Leu Tyr Asp Val 865 870 875 880

Gly Asp Ser Thr Ile Lys Asp Lys Ser Ala Leu Lys Leu Asn Phe Ala 885 890 895

Asp Arg Phe Asn Gly Ser Asn Glu Ala Lys Gln Thr Asp Asn Leu His
900 905 910

Leu Pro Ile Leu Pro Pro Leu Asn Gly Asp Asn Glu Leu Arg Lys Gln 915 920 925

Asn Ser Gln Glu Gly Asp Gln Ala His Pro Lys Ile Lys Ser Met Ile 930 935 940

Pro Glu Ser Gly Ser Ser Ser His Thr Glu Lys Glu Glu Glu Asn Glu 945 950 955 960

Glu Lys Glu Glu Lys Lys Pro Glu Gln His Lys Gln Glu Glu Asp Gln
965 970 975

Glu Lys Arg Glu Lys Val Val Asp Asp Met Glu Pro Pro Leu Asn Lys 980 985 990

Ser Val Gln Lys Ile Arg Glu Lys Asn Ala Gly Ser Gln Ala Lys Asp 995 1000 1005

His Ser Lys Asp His Leu Lys Glu His Lys Gln Asp Lys Asn Thr Ala 1010 1015 1020

Ile Gly Asn Gly Ser Phe Phe Arg Lys Phe Ser Lys Ser Ser Asp Lys

SEQL

1025 1030 1035 1040

Thr Met Glu Leu Tyr Ala Lys Ile Ser Ala Lys Gln Leu Phe Asn Gly
1045 1050 1055

Leu Glu Lys Leu Leu Arg Gly Trp Thr Gln Tyr Gly Leu Lys Asn Ile 1060 1065 1070

Lys Ser His Pro Asn Asn Leu Thr Leu Thr Gly Lys Leu Ser Ser Asp 1075 1080 1085

Asn Ile Phe Ser Leu Arg Ser Thr Leu Phe Glu Val Asn Ile Tyr Pro 1090 1095 1100

Arg Gly Lys Met Ser Val Val Gln Phe Lys Lys Val Ser Gly Ser Phe 1105 1110 1115 1120

Lys Ala Val Lys Lys Leu Val Asn Glu Val Glu Asn Val Leu Asn Lys 1125 1130 1135

Glu Gly Val Leu Gln Lys 1140

<210> 73

<211> 1844

<212> DNA

<213> Saccharomyces cerevisiae

<400> 73

gagttgaatc tatgcgtaac cattttgact accgcttcgt atgctttctt gcactttgtg 60 gagctactca tacaatagct tataatctgt gtagtcaaac tatatactag gcaaattact 120 agtaaatgca gtattcatat gtcctcaaac cgttttttgc agtatgttat tgatccatcg 180 aatcaaaatt tttcaccgcc aagagaaaaa agatcttact attttgtgct tatgaaaaaa 240 ttagtaaaag taaagctgga agcacataca agcgcagctg aatcacaaag gaaaattggc 300 acatecttgg ttgacacttg tetatttatt aaaaggtttt aaaaagtagt gttcageagt 360 tctatcatat ttttttattt aaaaaaaaat actgagagga agcgtattgt taagacacaa 420 tttagggtgg cgttaaataa acgagagagc ccaaaaatat aaccaagata aagaaaatca 480 atcataaagt gaattcaaaa atgtcatcgc aaaacctcaa tgataatcca aaaaatacat 540 cctcagcagc tgaagataag aagaaacaaa cttcatcctt aaagttggca ccaataccaa 600 ccacatcgcc atggaaatca tcttcgccag atagcaatac agtaattcct gtagaggaac 660 caagcaatac caaatggact ccaattacgc cgtctgttat aatctcaggt tccaaggaca 780 caaattcaaa gtcaggaaag aactctaaaa attctaagac taataaaaaa atgaaaaagc 840 gtggcaaata taataacgat atcaataaaa aggactttaa cggtcaaacc aacagtacat 900 cggaaataag taacgtttcc aatcttgaat ccaaaccttt agatgctaac gctaaagtaa 960 acatacattc aagctcagga gcaactgcca atgggaatat aaagaggata acaaacaaca 1020

```
acaattcaac caacggtaga caatcaagaa attatcaaaa caggaatggc aaaacaagat 1080 acaacaacaa tagtagacac agtcaggcgg ctaataatgc catctccttc ccaaataatt 1140 atcaggctag acctgaatat attcccaatg ccagccactg gttgaacaac aattcaagaa 1200 atagctataa acaactgtca tacttccgtc aacagcagta ttataataac atcaactatc 1260 aacaacaatt gcaaacacca tattattact caatggaacc tatttttaaa tctatcgaaa 1320 gtatcaaaaa ccaaattgaa ttctattta gtgaagagaa cttgaaaaca gatgaatttt 1380 taagatctaa attcaaaaaa gccaatgacg gatttatccc catgagtttg atagggaaat 1440 tttaccgtat ggttaattta tctcttggtg gagacccaaa tttaattttg gcatctatga 1500 gagaagtttt acaacataaa gaaacaaacc atttggaaat tgcccttgga agcatagaag 1560 gtgctcagaa gaacatggca gatgatttca atccattgga aaactattt attaggcgcg 1620 aaaattgggc tgaatacgct atggaaagta attttgatga aaatgatgac gaacatgaaa 1680 atccaaactt ctttcccagt aatggaactg ggaaaaagag tcagagctat gaccaaggtg 1800 aaattagcag gcagtttgaa caaaacttac aaaataaaga gcagtttgaa ttaa
```

<210> 74

<211> 447

<212> PRT

<213> Saccharomyces cerevisiae

<400> 74

Met Ser Ser Gln Asn Leu Asn Asp Asn Pro Lys Asn Thr Ser Ser Ala 1 5 10 15

Ala Glu Asp Lys Lys Gln Thr Ser Ser Leu Lys Leu Ala Pro Ile 20 25 30

Pro Thr Thr Ser Pro Trp Lys Ser Ser Ser Pro Asp Ser Asn Thr Val
35 40 45

Ile Pro Val Glu Glu Leu Arg Asp Ile Ser Lys Thr Ala Lys Pro Ser 50 55 60

Lys Asn Gly Ser Gly Ser Ile Lys Leu Thr Ser Asn Thr Lys Trp Thr 65 70 75 80

Pro Ile Thr Pro Ser Val Ile Ile Ser Gly Ser Lys Asp Thr Asn Ser 85 90 95

Lys Ser Gly Lys Asn Ser Lys Asn Ser Lys Thr Asn Lys Lys Met Lys
100 105 110

Lys Arg Gly Lys Tyr Asn Asn Asp Ile Asn Lys Lys Asp Phe Asn Gly
115 120 125

Gln Thr Asn Ser Thr Ser Glu Ile Ser Asn Val Ser Asn Leu Glu Ser 130 135 140

Lys	Pro	Leu	Asp	Ala	Asn	Ala	Lys	Val	Asn	Ile	His	Ser	Ser	Ser	GIA
145					150					155					160

- Ala Thr Ala Asn Gly Asn Ile Lys Arg Ile Thr Asn Asn Asn Asn Ser 165 170 175
- Thr Asn Gly Arg Gln Ser Arg Asn Tyr Gln Asn Arg Asn Gly Lys Thr 180 185 190
- Arg Tyr Asn Asn Asn Ser Arg His Ser Gln Ala Ala Asn Asn Ala Ile 195 200 205
- Ser Phe Pro Asn Asn Tyr Gln Ala Arg Pro Glu Tyr Ile Pro Asn Ala 210 215 220
- Ser His Trp Leu Asn Asn Asn Ser Arg Asn Ser Tyr Lys Gln Leu Ser 225 230 235 235
- Tyr Phe Arg Gln Gln Gln Tyr Tyr Asn Asn Ile Asn Tyr Gln Gln Gln 245 250 255
- Leu Gln Thr Pro Tyr Tyr Tyr Ser Met Glu Pro Ile Phe Lys Ser Ile 260 265 270
- Glu Ser Ile Lys Asn Gln Ile Glu Phe Tyr Phe Ser Glu Glu Asn Leu 275 280 285
- Lys Thr Asp Glu Phe Leu Arg Ser Lys Phe Lys Lys Ala Asn Asp Gly 290 295 300
- Phe Ile Pro Met Ser Leu Ile Gly Lys Phe Tyr Arg Met Val Asn Leu 305 310 315 320
- Ser Leu Gly Gly Asp Pro Asn Leu Ile Leu Ala Ser Met Arg Glu Val 325 330 335
- Leu Gln His Lys Glu Thr Asn His Leu Glu Ile Ala Leu Gly Ser Ile 340 345 350
- Glu Gly Ala Gln Lys Asn Met Ala Asp Asp Phe Asn Pro Leu Glu Asn 355 360 365
- Tyr Phe Ile Arg Arg Glu Asn Trp Ala Glu Tyr Ala Met Glu Ser Asn 370 375 380
- Phe Asp Glu Asn Asp Asp Glu Thr Glu Lys Tyr Asn Ile Glu Lys Leu 385 390 395 400



Leu Gly Pro Asn Asp Leu Asp Asn Tyr Ser Tyr Met Gly Tyr Pro Asn 405 410 415

Phe Phe Pro Ser Asn Glu Asn Gly Lys Lys Ser Gln Ser Tyr Asp Gln
420 425 430

Gly Glu Ile Ser Arg Gln Phe Glu Gln Asn Leu Gln Ile Asn Asp 435 440 445

<210> 75

<211> 2054

<212> DNA

<213> Saccharomyces cerevisiae ·

## <400> 75

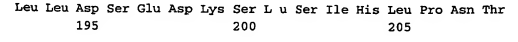
gcagcagggg caagatgaca accetgttee tgtteetgtt ecagtagaat etgagaegge 60 ttttgtgcca tcagcattca cagcaccacc ggtgccaaca aagaagaaat caaaaaataa 120 aaagggtacc cagccattgg caatggatga ctattttaat gaaggcagag ataagtcatc 180 taccgctgcg aagtcagcag aatctgacat cctcgcccca ccaccacaaa aacagtcatc 240 ctctgattaa ctttcctggt tagtcttttg gttttgtttc atagcaaaat taaatatata 300 tatataagct tgctttccct tcaaaacacg taaacgatag ttggcaatgt acgaaaagta 360 ccgagacttt ttttcaaagg cacgcgtgtc cttttttgtt aagacaatag atattttagc 420 attcagaaag tttcaatttc caagacttga cgtttcaatt atatggcaat ctcccaacaa 480 gcacccgctc atataatacc atgcaagtga ccacaagatt tatatctgcg atagtctcgt 540 tttgcctgtt tgcttctttc acgttggctg aaaacagcgc aagagctacg ccgggatcag 600 atttactcgt tctaacagag aagaaattta aatcattcat cgaatctcat ccgttagtcc 660 tcgtcgagtt ttttgctcca tggtgtttgc attctcagat cttacgccct cacttagaag 720 aggccgcctc tattttaaag gagcataacg tcccagttgt tcaaattgat tgtgaggcta 780 acagtatggt ttgcctgcaa caaactataa atacctaccc aaccttgaaa atctttaaaa 840 atggtcgtat ttttgatggt caagtctatc gcggtgtcaa gatcaccgat gaaatcactc 900 agtacatgat tcagctatac gaggettetg teatttattt aaatteegaa gatgaaatee 960 aaccatactt ggaaaatgca actttaccag tagtaataaa cagaggcttg acaggcttga 1020 atgaaacgta tcaagaagtc gcactggacc ttgctgagga ttacgtcttt ttatcccttc 1080 tagattcaga agataagtca ttatcaatcc acttgccaaa cactacagaa ccaattctgt 1140 ttgatggaaa tgtagactct ttggtcggaa attccgttgc tctaactcag tggttaaaag 1200 tggtaatttt accttacttt accgacatcg aacctgatct cttccccaag tacatttcta 1260 gcaatttgcc gttggcttac ttcttttata cttctgagga agaattggaa gattacactg 1320 atcttttcac gcagttaggt aaggaaaatc gtggccaaat aaatttcatt gcattaaact 1380 ctacaatgtt cccacaccac gttagattcc taaatatgag agaacagttc ccattatttg 1440 ctatccataa tatgatcaat aatctgaaat atggtttacc acaactacca gaagaagagt 1500 acgcgaaatt agaaaaacca caaccactag acagagatat gatcgttcag ttggtaaaag 1560 attaccgtga aggtactgcc aagccaattg ttaagtcaga agagattcca aaagaacaaa 1620 agtccaatgt ttataaaata gttgggaaga cacatgacga cattgttcat gatgatgaca 1680 aggatgteet tgteaaatat taegegacat ggtgtattea tagtaaaagg tttgegeeta 1740 tttacgaaga aattgcaaat gtcttagcat ctgatgaatc tgttcgcgat aaaatcttga 1800

tcgccgaagt agattcaggg gcaaatgata tcttaagttt tcctgtgaca ggatatccaa 1860 ccattgcttt gtatcctgcc ggaaataact ctaagcctat tatcttcaat aaaattagaa 1920 atttggaaga tgttttcgaa tttatcaagg aatcaggtac acatcacatt gacggccagg 1980 caatttatga taaattgcac caggccaagg attctgaagt gtctactgaa gataccgtac 2040 atgatgaatt ataa <210> 76 <211> 517 <212> PRT <213> Saccharomyces cerevisiae Met Gln Val Thr Thr Arg Phe Ile Ser Ala Ile Val Ser Phe Cys Leu 10 5 1 Phe Ala Ser Phe Thr Leu Ala Glu Asn Ser Ala Arg Ala Thr Pro Gly 25 30 20 Ser Asp Leu Leu Val Leu Thr Glu Lys Lys Phe Lys Ser Phe Ile Glu 40 Ser His Pro Leu Val Leu Val Glu Phe Phe Ala Pro Trp Cys Leu His 60 55 Ser Gln Ile Leu Arg Pro His Leu Glu Glu Ala Ala Ser Ile Leu Lys 75 80 70 Glu His Asn Val Pro Val Val Gln Ile Asp Cys Glu Ala Asn Ser Met 90 85 Val Cys Leu Gln Gln Thr Ile Asn Thr Tyr Pro Thr Leu Lys Ile Phe 100 Lys Asn Gly Arg Ile Phe Asp Gly Gln Val Tyr Arg Gly Val Lys Ile 120 115 Thr Asp Glu Ile Thr Gln Tyr Met Ile Gln Leu Tyr Glu Ala Ser Val 130 135

Ile Tyr Leu Asn Ser Glu Asp Glu Ile Gln Pro Tyr Leu Glu Asn Ala 145 150 160

Thr Leu Pro Val Val Ile Asn Arg Gly Leu Thr Gly Leu Asn Glu Thr 170 165

Tyr Gln Glu Val Ala Leu Asp Leu Ala Glu Asp Tyr Val Phe Leu Ser 190 180 185



- Thr Glu Pro Ile Leu Phe Asp Gly Asn Val Asp Ser Leu Val Gly Asn 210 215 220
- Ser Val Ala Leu Thr Gln Trp Leu Lys Val Val Ile Leu Pro Tyr Phe 225 230 235 240
- Thr Asp Ile Glu Pro Asp Leu Phe Pro Lys Tyr Ile Ser Ser Asn Leu 245 250 255
- Pro Leu Ala Tyr Phe Phe Tyr Thr Ser Glu Glu Glu Leu Glu Asp Tyr 260 265 270
- Thr Asp Leu Phe Thr Gln Leu Gly Lys Glu Asn Arg Gly Gln Ile Asn 275 280 285
- Phe Ile Ala Leu Asn Ser Thr Met Phe Pro His His Val Arg Phe Leu 290 295 300
- Asn Met Arg Glu Gln Phe Pro Leu Phe Ala Ile His Asn Met Ile Asn 305 310 315 320
- Asn Leu Lys Tyr Gly Leu Pro Gln Leu Pro Glu Glu Glu Tyr Ala Lys 325 330 335
- Leu Glu Lys Pro Gln Pro Leu Asp Arg Asp Met Ile Val Gln Leu Val
  340 345 350
- Lys Asp Tyr Arg Glu Gly Thr Ala Lys Pro Ile Val Lys Ser Glu Glu 355 360 365
- Ile Pro Lys Glu Gln Lys Ser Asn Val Tyr Lys Ile Val Gly Lys Thr 370 375 380
- His Asp Asp Ile Val His Asp Asp Lys Asp Val Leu Val Lys Tyr 385 390 395 400
- Tyr Ala Thr Trp Cys Ile His Ser Lys Arg Phe Ala Pro Ile Tyr Glu
  405 410 415
- Glu Ile Ala Asn Val Leu Ala Ser Asp Glu Ser Val Arg Asp Lys Ile 420 425 430
- Leu Ile Ala Glu Val Asp Ser Gly Ala Asn Asp Ile Leu Ser Phe Pro 435 440 445



```
Val Thr Gly Tyr Pro Thr Ile Ala Leu Tyr Pro Ala Gly Asn Asn Ser
                                            460
    450
                        455
Lys Pro Ile Il Phe Asn Lys Ile Arg Asn Leu Glu Asp Val Phe Glu
465
                    470
                                        475
Phe Ile Lys Glu Ser Gly Thr His His Ile Asp Gly Gln Ala Ile Tyr
                                    490
                485
Asp Lys Leu His Gln Ala Lys Asp Ser Glu Val Ser Thr Glu Asp Thr
                                505
                                                     510
Val His Asp Glu Leu
        515
<210> 77
<211> 908
<212> DNA
<213> Saccharomyces cerevisiae
<400> 77
ggggcaaatg atatcttaag ttttcctgtg acaggatatc caaccattgc tttgtatcct 60
gccggaaata actctaagcc tattatcttc aataaaatta gaaatttgga agatgttttc 120
gaatttatca aggaatcagg tacacatcac attgacggcc aggcaattta tgataaattg 180
caccaggeca aggattetga agtgtetaet gaagataeeg tacatgatga attataatea 240
ataaataaag catatataat gcacattttt aacatctgat tactcgcatc gtttctggaa 300
gaaaatagct aatattegtt atttatggca teaegattat teteaecagt taccegttta 360
tgctcttgaa gagatttagc attactgcca gcgcatcttc aaatacaggt ttatatgaga 420
cccattacta taaccctaag aagagaaaaa ggagtgcttt cgttttcaat cacattctag 480
tttacagtaa ttgagtctcg atgatgttta atatttacct tttcgtcact ttttttcca 540
ccattcttgc aggttccctg tcagatttgg aaatcggtat tatcaagaga ataccggtag 600
aagattgctt aattaaggca atgccaggtg ataaagttaa ggttcattat acaggatctt 660
tattagaatc gggaactgta tttgactcaa gttattcaag aggctctcct atcgcttttg 720
aacttggcgt tggcagagta attaaaggtt gggatcaagg tgttgccggc atgtgcgttg 780
gcgaaaaaag aaagctgcaa attccaagtt ctttggccta cggagaaaga ggtgtcccag 840
gcgtcattcc tccaagtgct gatttggtgt ttgatgtcga attggtagac gtgaaatcag 900
                                                                   908
ccgcctag
<210> 78
<211> 135
<212> PRT
<213> Saccharomyces cerevisiae
```

<400> 78

02-07-1999

Met Met Phe Asn Ile Tyr Leu Phe Val Thr Phe Phe Ser Thr Ile Leu 10

Ala Gly Ser Leu Ser Asp Leu Glu Ile Gly Ile Ile Lys Arg Ile Pro 25

Val Glu Asp Cys Leu Ile Lys Ala Met Pro Gly Asp Lys Val Lys Val 35 40

His Tyr Thr Gly Ser Leu Leu Glu Ser Gly Thr Val Phe Asp Ser Ser 50 55

Tyr Ser Arg Gly Ser Pro Ile Ala Phe Glu Leu Gly Val Gly Arg Val 70 75

Ile Lys Gly Trp Asp Gln Gly Val Ala Gly Met Cys Val Gly Glu Lys 85 90

Arg Lys Leu Gln Ile Pro Ser Ser Leu Ala Tyr Gly Glu Arg Gly Val 100 105 110

Pro Gly Val Ile Pro Pro Ser Ala Asp Leu Val Phe Asp Val Glu Leu 120 125

Val Asp Val Lys Ser Ala Ala 130

<210> 79

<211> 1103

<212> DNA

<213> Saccharomyces cerevisiae

<400> 79

cgagettgec ageattecaa tgeecategt attgtactea atggtgaaac ttaccaccat 60 attgttaaaa gctattattt atgattttta cttaccaaac attgaaagga gactgaacac 120 accacgataa aacgtatgtt ctgataccca gcaaatgaga aagtagaagg agaaacctaa 180 aaacctgtgg cggtcaagac tgctagcaag atggatattt agatcataaa ttctaacgca 240 tgcatcctga gcaatgaaat catgcaacct aacatttgag gttaatatat gagaattaac 300 tagcgttgag gaggtactgc aatttaaaag accgaagaat tatcgatgca aggaaaaatg 360 ggtctaggat gaatacgagc aattgaaata catttggaat accttgtgaa aatatcacat 420 actttcgcct tctatctcga tgcgttatta caagaaaata gttttactaa caaattaaca 480 aaaattaaaa tagtgtaaaa atgggtattt ctcgtgattc tcgtcacaaa agatcagcca 540 ctggtgctaa acgtgctcaa ttcagaaaga agagaaagtt cgaattaggt cgtcaaccag 600 ccaacacaaa aattggtgct aagagaatcc actctgtcag aactagaggt ggtaacaaga 660 aatacagage tetaagaatt gaaaceggta acttttettg ggettetgaa ggtateteca 720 agaagaccag aattgctggt gttgtttacc atccatccaa caatgaattg gttagaacta 780 acactttgac caaggctgcc attgtccaaa ttgatgctac tccattcaga caatggttcg 840 aagctcacta cggtcaaacc ttgggtaaga agaagaacgt caaggaagaa gaaactgttg 900 ccaagagcaa gaacgctgaa agaaagtggg ctgctagagc tgcttctgcc aagatcgaat 960 cttccgttga atctcaattc agcgccggta gattatacgc ttgtatctct tccagaccag 1020 gtcaatccgg tagatgtgat ggttacatct tggaaggtga agaattagct ttctacctaa 1080 gaagattgac tgctaagaaa tag 1103

<210> 80

<211> 200

<212> PRT

<213> Saccharomyces cerevisiae

<400> 80

Met Gly Ile Ser Arg Asp Ser Arg His Lys Arg Ser Ala Thr Gly Ala 1 5 10 15

Lys Arg Ala Gln Phe Arg Lys Lys Arg Lys Phe Glu Leu Gly Arg Gln 20 25 30

Pro Ala Asn Thr Lys Ile Gly Ala Lys Arg Ile His Ser Val Arg Thr
35 40 45

Arg Gly Gly Asn Lys Lys Tyr Arg Ala Leu Arg Ile Glu Thr Gly Asn 50 55 60

Phe Ser Trp Ala Ser Glu Gly Ile Ser Lys Lys Thr Arg Ile Ala Gly 65 70 75 80

Val Val Tyr His Pro Ser Asn Asn Glu Leu Val Arg Thr Asn Thr Leu
85 90 95

Thr Lys Ala Ala Ile Val Gln Ile Asp Ala Thr Pro Phe Arg Gln Trp
100 105 110

Phe Glu Ala His Tyr Gly Gln Thr Leu Gly Lys Lys Lys Asn Val Lys 115 120 125

Glu Glu Glu Thr Val Ala Lys Ser Lys Asn Ala Glu Arg Lys Trp Ala 130 135 140

Ala Arg Ala Ala Ser Ala Lys Ile Glu Ser Ser Val Glu Ser Gln Phe 145 150 155 160

Ser Ala Gly Arg Leu Tyr Ala Cys Ile Ser Ser Arg Pro Gly Gln Ser 165 170 175

Gly Arg Cys Asp Gly Tyr Ile Leu Glu Gly Glu Leu Ala Phe Tyr

185

190

Leu Arg Arg Leu Thr Ala Lys Lys 195 200

<210> 81

<211> 1265

<212> DNA

<213> Saccharomyces cerevisiae

<400> 81

```
accetatatg ggaggacaac tegeceactg tttgggtegt gttagegata aggteegaag 60
tagaagcgga aagagaagga gctgaagagg ttaattcatc gatggaaggc aactgggttt 120
ggatttccga gacatcgttg gcatttgggc ccgtcgaatt aaatcttttg gcctgaaaag 180
agatecatga eggatggge eggggeaata etatggtteg ageggtggee agtetggaag 240
aggcagcaaa ccttgacgtg acgagtcgag aggtgagttt gaacatcgtc ggggaggtta 300
ttctgtggct ccgcttgtac gtgaacagat acgtatagag ggcgagccac tggttaaatt 360
tttcatggct cggattactt ccgtactgct ggctaaaatc gaaatctcgg cctgctgaga 420
gtgttttgag caatcaaggg aacatctgaa cgtggaagag cagacgaggc attagctcga 480
acataagaac ggaacacgtc atgttgacta tcacgaaaag actggtgacc accgatgtgc 540
ggtcgcgaat actgttaagc agtttaaacg ggaaaatgtc cgatgcactg gcgctgctgc 600
gtcagcagca gcagaccagc gtggatgtgg agctgctgca cacgatgcta gcgcgagccg 660
ctgegettge ceatgeegae actatageat acatgtggta teageatgtg atgecaegee 720
ggttgccagt agagggccgc ctgctatgtg aaatggctgg cgtagcattg taccaggaca 780
ggctcttctt acccgcgcag ttcctccagc actaccaggc gatgaatcgc gatcgtcgca 840
ccagcccaga agatgaactg attgagtatg agcttagacg gattaaagtc gaagcgtttg 900
cgcgtggcac aatgcactcc acggcgctca gggaaaagtg gaaggtattc ttgcaggaga 960
tggatacgct accagggcag ccgccattaa ggctgcgcga cttcccgcaa atgaccaagg 1020
ctatgggcat agcattgatg cagcaagatg agcaagcagc tgccctggcg ttgtttggac 1080
gacagcccct agtgataaag aacgaatggt cactaccgct actactggct ggtgtccttt 1140
ggcatgttcc cggcccagcg caggcgcgac gtgtgctggc ggagttccgt caaagttatc 1200
gcgggctgcc gctgctggat gccgaactag tgataaagag aagaggattt gaaatcaaca 1260
                                                                  1265
cataa
```

<210> 82

<211> 254

<212> PRT

<213> Saccharomyces cerevisiae

<400> 82

Met Leu Thr Ile Thr Lys Arg Leu Val Thr Thr Asp Val Arg Ser Arg

1 5 10 15

Ile Leu Leu Ser Ser Leu Asn Gly Lys Met Ser Asp Ala Leu Ala Leu 20 25 30

Leu Arg Gln Gln Gln Gln Thr Ser Val Asp Val Glu Leu Leu His Thr
35 40 45

Met Leu Ala Arg Ala Ala Leu Ala His Ala Asp Thr Ile Ala Tyr 50 55 60

Met Trp Tyr Gln His Val Met Pro Arg Arg Leu Pro Val Glu Gly Arg 65 70 75 80

Leu Leu Cys Glu Met Ala Gly Val Ala Leu Tyr Gln Asp Arg Leu Phe 85 90 95

Leu Pro Ala Gln Phe Leu Gln His Tyr Gln Ala Met Asn Arg Asp Arg 100 105 110

Arg Thr Ser Pro Glu Asp Glu Leu Ile Glu Tyr Glu Leu Arg Arg Ile 115 120 125

Lys Val Glu Ala Phe Ala Arg Gly Thr Met His Ser Thr Ala Leu Arg 130 135 140

Glu Lys Trp Lys Val Phe Leu Gln Glu Met Asp Thr Leu Pro Gly Gln 145 150 155 160

Pro Pro Leu Arg Leu Arg Asp Phe Pro Gln Met Thr Lys Ala Met Gly 165 170 175

Ile Ala Leu Met Gln Gln Asp Glu Gln Ala Ala Ala Leu Ala Leu Phe 180 185 190

Gly Arg Gln Pro Leu Val Ile Lys Asn Glu Trp Ser Leu Pro Leu Leu 195 200 205

Leu Ala Gly Val Leu Trp His Val Pro Gly Pro Ala Gln Ala Arg Arg

Val Leu Ala Glu Phe Arg Gln Ser Tyr Arg Gly Leu Pro Leu Leu Asp 225 230 235 240

Ala Glu Leu Val Ile Lys Arg Arg Gly Phe Glu Ile Asn Thr 245 250

<210> 83

<211> 830

<212> DNA

SEQL

## <213> Saccharomyces cerevisiae

<400> 83

ggactagaag ccaaaagcca gaggcggtaa aaatagcaag actagaatat tgctggcatc 60 tgttaagggg atatgttgca acttgcaggg ggcggcacaa aataacatag aaacgtagta 120 aagaggggaa aaggaaaagg aaaaggaaaa ggaaggaaaa aaacccattg acgtagaaat 180 tgaaagaagg aaaggtatac gcaagcatta atacaaccca caaacacaga ccagaagcac 240 tctagacgga gagtaactag atctacagcc cctggaaaat cgtttggtca actttgaggt 300 teeggtegte eccetettga tetgaaaggt etttetetaa atetatatta aaacgtataa 360 ataggacggt gaattgcgtt ctacttcctc aattgcgttt gatcttattt aatctctctc 420 taatatatag aaaaaaaaac catctgatta ttcgataatc tcaaacaaac aactcaaaac 480 aaaaaaaact aaatacaaca atgtctgacg caggtagaaa aggattcggt gaaaaagctt 540 ctgaagcttt gaagccagac tctcaaaagt catacgctga acaaggtaag gaatacatca 600 ctgacaaggc cgacaaggtc gctggtaagg ttcaaccaga agacaacaag ggtgtcttcc 660 aaggtgtcca cgactctgcc gaaaaaggca aggataacgc tgaaggtcaa ggtgaatctt 720 tggcagacca agctagagat tacatgggag ccgccaagtc caagttgaac gatgccgtcg 780 aatatgtttc cggtcgtgtc cacggtgaag aagacccaac caagaagtaa 830

<210> 84

<211> 109

<212> PRT

<213> Saccharomyces cerevisiae

<400> 84

Met Ser Asp Ala Gly Arg Lys Gly Phe Gly Glu Lys Ala Ser Glu Ala 1 10

Leu Lys Pro Asp Ser Gln Lys Ser Tyr Ala Glu Gln Gly Lys Glu Tyr 20 25

Ile Thr Asp Lys Ala Asp Lys Val Ala Gly Lys Val Gln Pro Glu Asp 40

Asn Lys Gly Val Phe Gln Gly Val His Asp Ser Ala Glu Lys Gly Lys 50 55 60

Asp Asn Ala Glu Gly Gln Gly Glu Ser Leu Ala Asp Gln Ala Arg Asp 65 70 75 80

Tyr Met Gly Ala Ala Lys Ser Lys Leu Asn Asp Ala Val Glu Tyr Val 85 90

Ser Gly Arg Val His Gly Glu Glu Asp Pro Thr Lys Lys 100 105

<210> 85

```
<211> 995
<212> DNA
<213> Saccharomyces cerevisiae
<400> 85
gaccttcage gttatecttg cetttttegg cagagtegtg gacacettgg aagacaceet 60
tgttgtcttc tggttgaacc ttaccagcga ccttgtcggc cttgtcagtg atgtattcct 120
taccttgttc agcgtatgac ttttgagagt ctggcttcaa agcttcagaa gctttttcac 180
cgaatcettt tetacetgeg teagacattg ttgtatttag ttttttttgt tttgagttgt 240
ataagatcaa acgcaattga ggaagtagaa cgcaattcac cgtcctattt atacgtttta 360
atatagattt agagaaagac ctttcagatc aagaggggga cgaccggaac ctcaaagttg 420
accaaacgat tttccagggg ctgtagatct agttactctc cgtctagagt gcttctggtc 480
tgtgtttgtg ggttgtatta atgcttgcgt atacctttcc ttctttcaat ttctacgtca 540
atgggttttt ttccttcctt ttccttttcc ttttcctttt cccctcttta ctacgtttct 600
atgttatttt gtgccgcccc ctgcaagttg caacatatcc ccttaacaga tgccagcaat 660
attetagtet tgetattttt accgeetetg gettttgget tetagteett gteccaagag 720
ccaagggccc gtcaacacgt cgtcattgct accgccagct ggcacccaca catcaccgac 780
cctttttttc cattttcggc tgggccgtta gtgggatccg cccgctcccg gagattttca 840
cttggatttg cgcgtcccct ttttttcttc attctctgac tccccctacc ttctcccact 900
tttctgtgta tcaagaggaa aagaaggaga aaaggagaac tccgaaaaat accgaacaag 960
                                                                995
agggtaacag aatgtgcatt tggatgagcg ggtaa
<210> 86
<211> 164
<212> PRT
<213> Saccharomyces cerevisiae
<400> 86
Met Leu Ala Tyr Thr Phe Pro Ser Phe Asn Phe Tyr Val Asn Gly Phe
                                    10
Phe Ser Phe Leu Phe Leu Phe Leu Phe Leu Phe Pro Ser Leu Leu Arg
                                25
Phe Tyr Val Ile Leu Cys Arg Pro Leu Gln Val Ala Thr Tyr Pro Leu
        35
                            40
Asn Arg Cys Gln Gln Tyr Ser Ser Leu Ala Ile Phe Thr Ala Ser Gly
                        55
                                            60
    50
Phe Trp Leu Leu Val Leu Val Pro Arg Ala Lys Gly Pro Ser Thr Arg
65
                    70
                                        75
                                                           80
Arg His Cys Tyr Arg Gln Leu Ala Pro Thr His His Arg Pro Phe Phe
                85
                                    90
```

```
Ser Ile Phe Gly Trp Ala Val Ser Gly Ile Arg Pro Leu Pro Glu Ile
100 Phe Thr Trp Ile Cys Ala Ser Pro Phe Phe Leu His Ser Leu Thr Pro
115 Phe Thr Phe Ser His Phe Ser Val Tyr Gln Glu Glu Lys Lys Glu Lys
130 Arg Arg Thr Pro Lys Asn Thr Glu Gln Glu Gly Asn Arg Met Cys Ile
```

Trp Met Ser Gly

145

```
<210> 87
<211> 2702
<212> DNA
<213> Saccharomyces cerevisiae
```

150

## <400> 87

```
ctgccttccg tacgtcacaa cgcatacaca atatagttta tataaactcc tgcgtataat 60
ctatectttt geataatatt ttegteacea ttatettgtt geggtgtttt tatttaaaac 120
aatagaactc gcctaaaggg gaaattttcg atataaaaat tcaaaaaaat ggctttcatg 180
gatcgagtat ttgtttgtcg aaaaaggatc actggagtga cgttactacg ctacgaagcc 240
teetteegge ttagecetgt tgattaegaa tttggateag ttggtatttg atteetegga 300
agagttaact tcacgcgggt aaatcacttg tgatgcggta tattctctat acggctaata 360
gatgaatcag ggtgttttaa agtgcgtata aacccttttg ctattttcgt ttatataatt 420
ggcgttgata aagagccaat atctattgtt gctacataga ggcagctctc ttagcaaaat 480
aaaaatacaa aaagttcgac atgggcttca gtagcggtaa atcaactaag aaaaagcctc 540
tgcttttcga tatcagactt aaaaatgttg acaacgatgt aatactcctc aaaggtcctc 600
caaacgaggc cccctcggtg cttttatctg gttgcatcgt tttatcgatt aacgaaccca 660
tgcagatcaa aagcatatca ttgagacttt atgggaagat acaaatagac gtaccattag 720
agaggececa ggaegetagt tettegtegt tgtetteate geegecaaag atcagaaagt 780
acaacaaagt tttttataat tacgcatggg ataatgttaa cctcaaggag tatctgagtg 840
gtttaagagg gcaatctggc cttgcgggca gtagctcatc aagtaatatc ttgggcactc 900
gccaaagagc tcagtccaca agttccttga agtctttaaa ggggtcctcc tcaccctctt 960
catgtacttt agataagggc aactacgatt ttccctttag tgctattttg cctggttcgt 1020
taccagagag cgtagaatct ttgccaaatt gcttcgtgac atatagcatg gaatccgtta 1080
ttgaacgcag caaaaattat agtgatttga tctgtaggaa aaatattaga gttctgagaa 1140
ccatttcacc cgcagcagtg gagttatcag aaactgtttg tgtagataac tcatggcccg 1200
acaaagtgga ttattctatt tcagtaccca acaaagccgt agctattggt tcagccaccc 1260
ctataaatat ttccattgta cctctttcga aaggtttgaa attgggctca atcaaagtcg 1320
tattatttga gaattatcaa tattgtgacc ccttccctcc agtaatttct gaaaataggc 1380
aagtgacaga actaaatett gaagateeet tgaacgagte atetggagaa tttaatggta 1440
```

```
atggttgctt tgtaaataac ccctttttc agcctgatca ttcattccaa gacaagtggg 1500
agattgatac catcctgcaa atcccgaaca gcttatcaaa ctgtgtgcaa gattgtgatg 1560
tccgctctaa cattaaggtt cgccataagc tcaaattttt catcatccta attaacccag 1620
atggtcataa atctgagtta agagcgtcct taccgattca actttttatt tcaccatttg 1680
tggcactttc aataaaacca ttgtcatcct cgaatttgta ttcgcttttt agcaccacta 1740
accagaaaga cgaaaactca tcacaagaag aggaagagga atatctgttt tctagatcag 1800
catcagtcac agggttggaa ttattagcgg atatgcgtag cggtggctct gttcctacca 1860
tttcagactt gatgacgccc ccaaattatg aaatgcacgt atatgatcgt ctttatagcg 1920
gttctttcac tcgcacggct gtggaaacgt ctggaacatg tactcctttg ggaagcgaat 1980
gttcgactgt cgaggatcag caacaggatt tagaagattt acgtatacgg ttgacaaaaa 2040
ttagaaatca acgtgacaat ctagggctac caccgtctgc ctcgtctgct gccgcttcca 2100
gatcgctatc tccattacta aacgttccag caccagagga tggcacggag agaatcttac 2160
ctcagagtgc tcttggtccc aatagtggct ctgtgccagg agtacatagt aacgtatcac 2220
etgttttact ttcaagatcc ccagccccaa gcgtgtcagc ccatgaagtg ttaccagtgc 2280
cctcgggctt aaattatcca gagactcaaa acctgaacaa ggttccatcg tatggcaagg 2340
caatgaaata tgatatcatt ggtgaggacc ttcctccttc ctacccttgt gcgatacaaa 2400
atgtgcaacc aagaaaaccc agtagggtac attccaggaa ctcttcgaca acattgtcat 2460
cttctatacc aactagcttt cattcctcta gttttatgag tagcactgct tcccctattt 2520
ccataattaa tggctctaga agtagttcta gtggggtatc tcttaataca cttaatgagt 2580
taacttegaa aacttegaat aacceateea gtaatagtat gaaaaggtea ceaacaagae 2640
ggagggctac ttctttagct gggtttatgg gaggttttct atcaaagggt aacaaacgat 2700
                                                                  2702
ag
```

<210> 88

<211> 733

<212> PRT

<213> Saccharomyces cerevisiae

<400> 88

Met Gly Phe Ser Ser Gly Lys Ser Thr Lys Lys Lys Pro Leu Leu Phe 1 5 10 15

Asp Ile Arg Leu Lys Asn Val Asp Asn Asp Val Ile Leu Leu Lys Gly
20 25 30

Pro Pro Asn Glu Ala Pro Ser Val Leu Leu Ser Gly Cys Ile Val Leu
35 40 45

Ser Ile Asn Glu Pro Met Gln Ile Lys Ser Ile Ser Leu Arg Leu Tyr
50 55 60

Gly Lys Ile Gln Ile Asp Val Pro Leu Glu Arg Pro Gln Asp Ala Ser 65 70 75 80

Ser Ser Ser Leu Ser Ser Pro Pro Lys Ile Arg Lys Tyr Asn Lys 85 90 95

Val	Phe	Tyr	Asn	Tyr	Ala	Trp	Asp	Asn	Val	Asn	Leu	Lys	Glu	Tyr	Leu
			100					105					110		

Ser Gly Leu Arg Gly Gln Ser Gly Leu Ala Gly Ser Ser Ser Ser Ser 115 120 125

Asn Ile Leu Gly Thr Arg Gln Arg Ala Gln Ser Thr Ser Ser Leu Lys 130 135 140

Ser Leu Lys Gly Ser Ser Ser Pro Ser Ser Cys Thr Leu Asp Lys Gly 145 150 155 160

Asn Tyr Asp Phe Pro Phe Ser Ala Ile Leu Pro Gly Ser Leu Pro Glu 165 170 175

Ser Val Glu Ser Leu Pro Asn Cys Phe Val Thr Tyr Ser Met Glu Ser 180 185 190

Val Ile Glu Arg Ser Lys Asn Tyr Ser Asp Leu Ile Cys Arg Lys Asn 195 200 205

Ile Arg Val Leu Arg Thr Ile Ser Pro Ala Ala Val Glu Leu Ser Glu 210 215 220

Thr Val Cys Val Asp Asn Ser Trp Pro Asp Lys Val Asp Tyr Ser Ile 225 230 235 240

Ser Val Pro Asn Lys Ala Val Ala Ile Gly Ser Ala Thr Pro Ile Asn 245 250 255

Ile Ser Ile Val Pro Leu Ser Lys Gly Leu Lys Leu Gly Ser Ile Lys
260 265 270

Val Val Leu Phe Glu Asn Tyr Gln Tyr Cys Asp Pro Phe Pro Pro Val 275 280 285

Ile Ser Glu Asn Arg Gln Val Thr Glu Leu Asn Leu Glu Asp Pro Leu 290 295 300

Asn Glu Ser Ser Gly Glu Phe Asn Gly Asn Gly Cys Phe Val Asn Asn 305 310 315 320

Pro Phe Phe Gln Pro Asp His Ser Phe Gln Asp Lys Trp Glu Ile Asp 325 330 335

Thr Ile Leu Gln Ile Pro Asn Ser Leu Ser Asn Cys Val Gln Asp Cys 340 345 350

- Asp Val Arg Ser Asn Ile Lys Val Arg His Lys Leu Lys Phe Phe Ile 355 360 365
- Ile Leu Ile Asn Pro Asp Gly His Lys Ser Glu Leu Arg Ala Ser Leu 370 375 380
- Pro Ile Gln Leu Phe Ile Ser Pro Phe Val Ala Leu Ser Ile Lys Pro 385 390 395 400
- Leu Ser Ser Ser Asn Leu Tyr Ser Leu Phe Ser Thr Thr Asn Gln Lys
  405 410 415
- Asp Glu Asn Ser Ser Gln Glu Glu Glu Glu Glu Tyr Leu Phe Ser Arg
  420 425 430
- Ser Ala Ser Val Thr Gly Leu Glu Leu Leu Ala Asp Met Arg Ser Gly
  435 440 445
- Gly Ser Val Pro Thr Ile Ser Asp Leu Met Thr Pro Pro Asn Tyr Glu 450 455 460
- Met His Val Tyr Asp Arg Leu Tyr Ser Gly Ser Phe Thr Arg Thr Ala 465 470 475 480
- Val Glu Thr Ser Gly Thr Cys Thr Pro Leu Gly Ser Glu Cys Ser Thr
  485 490 495
- Val Glu Asp Gln Gln Gln Asp Leu Glu Asp Leu Arg Ile Arg Leu Thr 500 505 510
- Lys Ile Arg Asn Gln Arg Asp Asn Leu Gly Leu Pro Pro Ser Ala Ser 515 520 525
- Ser Ala Ala Ala Ser Arg Ser Leu Ser Pro Leu Leu Asn Val Pro Ala 530 535 540
- Pro Glu Asp Gly Thr Glu Arg Ile Leu Pro Gln Ser Ala Leu Gly Pro 545 550 555
- Asn Ser Gly Ser Val Pro Gly Val His Ser Asn Val Ser Pro Val Leu
  565 570 575
- Leu Ser Arg Ser Pro Ala Pro Ser Val Ser Ala His Glu Val Leu Pro 580 585 590
- Val Pro Ser Gly Leu Asn Tyr Pro Glu Thr Gln Asn Leu Asn Lys Val 595 600 605

```
Pro Ser Tyr Gly Lys Ala Met Lys Tyr Asp Ile Ile Gly Glu Asp Leu
    610
                        615
                                             620
Pro Pro Ser Tyr Pro Cys Ala Ile Gln Asn Val Gln Pro Arg Lys Pro
625
                    630
                                        635
                                                             640
Ser Arg Val His Ser Arg Asn Ser Ser Thr Thr Leu Ser Ser Ser Ile
                645
                                     650
                                                         655
Pro Thr Ser Phe His Ser Ser Ser Phe Met Ser Ser Thr Ala Ser Pro
            660
                                 665
                                                     670
Ile Ser Ile Ile Asn Gly Ser Arg Ser Ser Ser Ser Gly Val Ser Leu
        675
                             680
                                                 685
Asn Thr Leu Asn Glu Leu Thr Ser Lys Thr Ser Asn Asn Pro Ser Ser
    690
                        695
                                             700
Asn Ser Met Lys Arg Ser Pro Thr Arg Arg Arg Ala Thr Ser Leu Ala
                    710
                                         715
Gly Phe Met Gly Gly Phe Leu Ser Lys Gly Asn Lys Arg
                725
                                    730
```

<210> 89 <211> 1259 <212> DNA <213> Saccharomyces cerevisiae

<400> 89

agacaataaa atatggaaaa taaaacgctt agcacagcag tcaatgaaat actctgatat 60 attttgactc tgtggagcat acagctacca ctaaatagga ttgtggggac aaatatggaa 120 tggatcaaaa cgttcttagg attgtggtgg taaaacttgt aaaatccaag ctgagatctt 180 aaatccagca aaccttcgcc catatttatt cttttataac agaagaagag actatattct 240 caaaacctcg tatatttata tacatatcct ccaaacaaac tcccaagttt cactttcctg 300 gatttacctt ggcattcctt ttccccatcc tcttataatg gtaatcgcga tccttaatta 360 tgatatcaca atagacgaag ggcacacacc aatttgccac cgtaaggata gagtaagtta 420 aatggctaac tcattataat cttcatgcta aatcatataa gggcagagac gaagcaaagc 480 gaaaaaaaca tattacaatc atgtcgggtg ctgctgctgc atctgctgct ggttatgaca 540 ggcacatcac tatcttttcc cccgagggtc gtttatatca agtagaatat gcctttaaag 600 cgactaatca aactaacata aactcactag cggtcagagg taaagattgt acagtggtga 660 taagtcagaa aaaggtccct gataaactgt tggatccaac tactgtttcg tatatttttt 720 gtatttcaag aacaattggt atggtagtga acggaccaat accggatgca agaaatgcgg 780 ccctaagagc caaggctgag gctgcagaat tccgttataa atatggttat gatatgccat 840 gcgatgtatt ggctaagaga atggctaacc tttcccaaat ctatactcaa agagcatata 900 tgagaccatt aggtgttata ctcacatttg tttcggtaga tgaagaattg ggtccctcca 960

tttacaaaac tgaccctgca ggttattacg ttggctacaa agctactgcg acaggaccaa 1020 aacaacagga gatcacaaca aacttagaaa accatttcaa aaagagtaaa atcgaccata 1080 ttaatgaaga atcatgggag aaggtggtag aatttgctat tactcatatg attgacgcac 1140 tgggtaccga attttcaaag aatgacttgg aagtcggtgt cgctacaaag gacaaattct 1200 ttaccttgag tgctgagaac atcgaagaaa ggctagtagc aattgctgaa caagattaa 1259

<210> 90

<211> 252

<212> PRT

<213> Saccharomyces cerevisiae

<400> 90

Met Ser Gly Ala Ala Ala Ala Ser Ala Ala Gly Tyr Asp Arg His Ile
1 5 10 15

Thr Ile Phe Ser Pro Glu Gly Arg Leu Tyr Gln Val Glu Tyr Ala Phe
20 25 30

Lys Ala Thr Asn Gln Thr Asn Ile Asn Ser Leu Ala Val Arg Gly Lys
35 40 45

Asp Cys Thr Val Val Ile Ser Gln Lys Lys Val Pro Asp Lys Leu Leu 50 55 60

Asp Pro Thr Thr Val Ser Tyr Ile Phe Cys Ile Ser Arg Thr Ile Gly
65 70 75 80

Met Val Val Asn Gly Pro Ile Pro Asp Ala Arg Asn Ala Ala Leu Arg 85 90 95

Ala Lys Ala Glu Ala Ala Glu Phe Arg Tyr Lys Tyr Gly Tyr Asp Met
100 105 110

Pro Cys Asp Val Leu Ala Lys Arg Met Ala Asn Leu Ser Gln Ile Tyr 115 120 125

Thr Gln Arg Ala Tyr Met Arg Pro Leu Gly Val Ile Leu Thr Phe Val 130 135 140

Ser Val Asp Glu Glu Leu Gly Pro Ser Ile Tyr Lys Thr Asp Pro Ala 145 150 155 160

Gly Tyr Tyr Val Gly Tyr Lys Ala Thr Ala Thr Gly Pro Lys Gln Gln
165 170 175

Glu Ile Thr Thr Asn Leu Glu Asn His Phe Lys Lys Ser Lys Ile Asp 180 185 190

```
His Ile Asn Glu Glu Ser Trp Glu Lys Val Val Glu Phe Ala Ile Thr
        195
                            200
His Met Ile Asp Ala Leu Gly Thr Glu Phe Ser Lys Asn Asp Leu Glu
    210
                        215
                                            220
Val Gly Val Ala Thr Lys Asp Lys Phe Phe Thr Leu Ser Ala Glu Asn
225
                    230
                                        235
                                                             240
Ile Glu Glu Arg Leu Val Ala Ile Ala Glu Gln Asp
                245
                                    250
<210> 91
<211> 968
<212> DNA
<213> Saccharomyces cerevisiae
<400> 91
ccaagttaag caagacacca agacaataac ttgagaggta tgtccctata tgatgatgat 60
tcataattaa tggcgaccga gggcagcgtt gtgctagaaa gcaatgtatt aatatatgag 120
agagetteag atageeaata caceggaatt tagaetttga caggattgtg gettgtteca 180
gtataactta taatatatgt caacetttga gcaggaaacc caatgaaaat gcttcattac 240
atatatacga tatggagaag catgaaaaat aacagcgtga gatgttatcc atgttgcaag 300
aacctagtaa aatgaattet gaatttgeaa aaagteeata ttteeaegtt eteetettee 360
tacaattttg cgaacgcctg aacaaccatg cggattacca ttatttatat tgacaagatg 420
gctacctatg aaaagcatag acttactaac atttttttt tcaaatatgt ttgaaaaacg 480
tggattaata tagtgataaa atgaaggttg aaatcgattc tttttcaggt gccaaaatct 540
acccaggcag aggtacettg tttgteegtg gtgacteeaa aatetteaga tteeaaaact 600
ccaaatctgc ctctttgttc aagcaaagaa agaacccaag aagaatcgct tggactgtct 660
tattcagaaa gcatcacaag aagggtatca ccgaagaagt tgctaagaag agatctagaa 720
aaaccgttaa ggcccaaaga ccaattaccg gtgcttcttt ggacttgatc aaggaaagaa 780
gatctttgaa gccagaagtt agaaaggcta acagagaaga aaaattgaag gccaacaaag 840
aaaagaagaa ggctgaaaag gctgctagaa aggctgaaaa ggctaagtct gctggtactc 900
aaagttctaa gttctccaag caacaagcta agggtgcttt ccaaaaggtt gctgctactt 960
ctcgttaa
<210> 92
<211> 155
<212> PRT
<213> Saccharomyces cerevisiae
<400> 92
```

Met Lys Val Glu Ile Asp Ser Phe Ser Gly Ala Lys Ile Tyr Pro Gly

```
Arg Gly Thr Leu Phe Val Arg Gly Asp Ser Lys Ile Phe Arg Phe Gln
                                 25
Asn Ser Lys Ser Ala Ser Leu Phe Lys Gln Arg Lys Asn Pro Arg Arg
                             40
Ile Ala Trp Thr Val Leu Phe Arg Lys His His Lys Lys Gly Ile Thr
                         55
Glu Glu Val Ala Lys Lys Arg Ser Arg Lys Thr Val Lys Ala Gln Arg
                     70
                                          75
 65
                                                              80
Pro Ile Thr Gly Ala Ser Leu Asp Leu Ile Lys Glu Arg Arg Ser Leu
                 85
                                      90
Lys Pro Glu Val Arg Lys Ala Asn Arg Glu Glu Lys Leu Lys Ala Asn
            100
                                105
Lys Glu Lys Lys Lys Ala Glu Lys Ala Ala Arg Lys Ala Glu Lys Ala
                            120
Lys Ser Ala Gly Thr Gln Ser Ser Lys Phe Ser Lys Gln Gln Ala Lys
                        135
Gly Ala Phe Gln Lys Val Ala Ala Thr Ser Arg
145
                    150
                                         155
<210> 93
<211> 764
<212> DNA
<213> Saccharomyces cerevisiae
<400> 93
```

ctcccatatt ttttattca caggctagac aatggtaatg gtgctttaga tgatttcagt 60 ggattacatt cgcacaatag aataaatgac taattttcaa gaatttgttt caacctgccg 120 gcattcagtt attataagtt attgtttgat ggtaaaactt cgccatcacc ggttgctgcg 180 tttattgcaa aattactgaa accatgtaat ttccgaatac ggtaattaca cgcagtttga 240 aatttgacat gcaagtttaa caaaaaaact tctgtccgca tctaaagctt aattaaggga 300 ctgtttcata ccttcagatc caatatactg aatactaggc aattgattc ccttgtaaat 360 atacaatata aatagttgta agaagaggtg atatgttaag tttccatttt gacgtattcc 420 tcattctaga atgattgtaa gctctcaaca gtcacttgtg tgccaaaata ttatacctac 480 ttcatacatt ttcaattaag atgcagttac ttcgctgttt ttcaatattt tctgttattg 540 cttcagttt agcacaggaa ctgacaacta tatgcgagca aatcccctca ccaactttag 600 gagtttttga atattacaaa tcagtaacgt ttgtcagtaa ttgcggttct cacccctcaa 720

SEQL

caactagcaa aggcagcccc ataaacacac agtatgtttt ttga

```
764
```

<210> 94

<211> 87

<212> PRT

<213> Saccharomyces cerevisiae

<400> 94

Met Gln Leu Leu Arg Cys Phe Ser Ile Phe Ser Val Ile Ala Ser Val 1 5 10 15

Leu Ala Gln Glu Leu Thr Thr Ile Cys Glu Gln Ile Pro Ser Pro Thr 20 25 30

Leu Glu Ser Thr Pro Tyr Ser Leu Ser Thr Thr Thr Ile Leu Ala Asn
35 40 45

Gly Lys Ala Met Gln Gly Val Phe Glu Tyr Tyr Lys Ser Val Thr Phe 50 55 60

Val Ser Asn Cys Gly Ser His Pro Ser Thr Thr Ser Lys Gly Ser Pro 65 70 75 80

Ile Asn Thr Gln Tyr Val Phe

85

<210> 95

<211> 1430

<212> DNA

<213> Saccharomyces cerevisiae

<400> 95

tcgggatggt gggttgttct gtaccccgc attcacaccg ctctgcctat tcatttatc 60 gttttcgtct tttcttcaa tgtattgata attaacgtgt ctatgtgatg ctattggtat 120 atagcctttc ttgtatatcc ctctaacaat tattacaagt gtttctgttt aaagcaagtg 180 ctagaagtga ctcggttctc ttaggaaatt ctaaacgcaa gatttctctt atggtgattg 240 taacaattat gagatacttc actagcccac cttaacttta cggaccttct tttgaaacga 300 tgcttcgaat gacaacgcct ttttgatata taatatccaa tttcattata gggaaatttt 360 caactcttac ccgcccact gtgctgatat gaccaagtga tcactcgatg atgggactac 420 gtattgaaaa atattgaatg aaaaattact caagcagcag aacattcaca gtgtagtcag 480 aaaagaacaa aagtaatgat gctgcagttc tagaaagtac ggtacatgtt aagaatctag 540 aaaagaacaa aagtaatgat gctgcagttc tagaaatctt acatgtcttg gataaagaat 600 tcgtcccac tgaaaagtta ctgagagaaa caaagttgg tgtggaagtc aacaagttta 660 aaaaaatccac taatgtagag atcagcaac tcgtgaagaa aatgattagc tcttggaaag 720 acgcaattaa taaaaataag cgttccaggc aagcacagca gcatcatcaa gatcatgcgc 780

caggcaatgc agaggacaag acaactgtag gtgagtccgt gaatggtgtt caacagccgg 840

```
cctcctccca gtcagatgcc atgaaacaag acaagtacgt cagcactaaa ccaagaaata 900
 gtaagaacga tggtgtggat acagctatat accaccacaa attacgtgat caggtactaa 960
 aagcactcta cgacgttttg gccaaggaaa gtgagcatcc acctcaatct attttgcata 1020
 ctgcaaaggc catagaaagt gaaatgaata aagttaacaa ctgtgacacc aacgaagccg 1080
cttacaaagc caggtatcgt ataatttatt caaacgtcat atcaaagaat aacccagatc 1140
tcaaacataa aattgccaac ggtgatataa cacctgaatt cttagctaca tgcgatgcca 1200
aggatctggc accagcgccc ttaaagcaaa agatagaaga aattgccaag caaaacttat 1260
acaacgcaca gggtgccacc atagaaaggt cagtcaccga tagatttaca tgtggtaaat 1320
gtaaagagaa gaaggtatct tactatcaat tgcaaacaag atctgcggat gaaccattga 1380
ccactttctg tacatgtgaa gcatgtggta acagatggaa attctcttag
<210> 96
<211> 309
<212> PRT
<213> Saccharomyces cerevisiae
<400> 96
Met Asp Ser Lys Glu Val Leu Val His Val Lys Asn Leu Glu Lys Asn
Lys Ser Asn Asp Ala Ala Val Leu Glu Ile Leu His Val Leu Asp Lys
                                  25
             20
Glu Phe Val Pro Thr Glu Lys Leu Leu Arg Glu Thr Lys Val Gly Val
         35
                              40
                                                  45
Glu Val Asn Lys Phe Lys Lys Ser Thr Asn Val Glu Ile Ser Lys Leu
     50
                          55
                                              60
Val Lys Lys Met Ile Ser Ser Trp Lys Asp Ala Ile Asn Lys Asn Lys
 65
                      70
                                          75
Arg Ser Arg Gln Ala Gln Gln His His Gln Asp His Ala Pro Gly Asn
Ala Glu Asp Lys Thr Thr Val Gly Glu Ser Val Asn Gly Val Gln Gln
            100
                                 105
                                                     110
Pro Ala Ser Ser Gln Ser Asp Ala Met Lys Gln Asp Lys Tyr Val Ser
        115
                             120
                                                 125
Thr Lys Pro Arg Asn Ser Lys Asn Asp Gly Val Asp Thr Ala Ile Tyr
    130
                         135
                                             140
His His Lys Leu Arg Asp Gln Val Leu Lys Ala Leu Tyr Asp Val Leu
145
                     150
                                         155
                                                              160
```

```
Ala Lys Glu Ser Glu His Pro Pro Gln Ser Ile Leu His Thr Ala Lys
                 165
                                     170
                                                          175
Ala Ile Glu Ser Glu Met Asn Lys Val Asn Asn Cys Asp Thr Asn Glu
             180
                                 185
                                                      190
Ala Ala Tyr Lys Ala Arg Tyr Arg Ile Ile Tyr Ser Asn Val Ile Ser
        195
                             200
                                                 205
Lys Asn Asn Pro Asp Leu Lys His Lys Ile Ala Asn Gly Asp Ile Thr
    210
                         215
                                             220
Pro Glu Phe Leu Ala Thr Cys Asp Ala Lys Asp Leu Ala Pro Ala Pro
225
                     230
                                         235
Leu Lys Gln Lys Ile Glu Glu Ile Ala Lys Gln Asn Leu Tyr Asn Ala
                 245
                                     250
Gln Gly Ala Thr Ile Glu Arg Ser Val Thr Asp Arg Phe Thr Cys Gly
            260
                                 265
Lys Cys Lys Glu Lys Lys Val Ser Tyr Tyr Gln Leu Gln Thr Arg Ser
        275
                             280
                                                 285
Ala Asp Glu Pro Leu Thr Thr Phe Cys Thr Cys Glu Ala Cys Gly Asn
    290
                         295
                                             300
Arg Trp Lys Phe Ser
305
<210> 97
<211> 929
<212> DNA
<213> Saccharomyces cerevisiae
<400> 97
tgttacgatc aacattcttg tccgtaatca gcatggtagt taaacccatt ctgtacatat 60
ggtcagatat actttcgcat gggtcaatac catggtcaac ccaaccagtt cttctttgcg 120
ttttcaaaag ttgtataata ttcaaaaacg ctaactgata atttgggtga ggtttcgaaa 180
```

gaattgccaa aatctccta ggtatattat cttctggctt ccagatgtta actgcagtca 240 ttttgtttt cctggtctt acctcgaata ggttttcttg ttttgttatt tgttttcttt 300 ttggacccta tgtttatata tggattttga aaatctttta aaaaagcgat aaaagcgttg 360 ggatctgcca ctggtaactt caaaatagac aagacgaaaa aacggtgaaa atgggtgata 420 gaaataatac agaagtagat gttgaattag attaaactga agatatataa tttattggaa 480 aatacataga gttttgttg atgcgcttaa gcgatcaatt caacaacacc accagcagct 540

```
ctgatttttt cttcagccaa cttggagacg aatctagctt tgacgataac tggaacattt 600
ggaattctac ccttacccaa gatcttaccg taaccggctg ccaaagtgtc aataactgga 660
gcagtttcct tagaagcaga tttcaagtat tggtctctct tgtcttctgg gatcaatgtc 720
cacaatttgt ccaagttcaa gactggcttc cagaaatgag cttgttgctt gtggaagtat 780
ctcataccaa ccttaccgaa ataacctgga tggtatttat ccatgttaat tctgtggtga 840
tgttgaccac cggccatacc tctaccaccg gggtgctttc tgtgcttacc gatacgacct 900
ttaccggctg tacaaaaaaa atattgtaa
<210> 98
<211> 142
<212> PRT
<213> Saccharomyces cerevisiae
<400> 98
Met Arg Leu Ser Asp Gln Phe Asn Asn Thr Thr Ser Ser Ser Asp Phe
                  5
                                     10
Phe Phe Ser Gln Leu Gly Asp Glu Ser Ser Phe Asp Asp Asn Trp Asn
                                                     30 -
             20
                                 25
Ile Trp Asn Ser Thr Leu Thr Gln Asp Leu Thr Val Thr Gly Cys Gln
                             40
Ser Val Asn Asn Trp Ser Ser Phe Leu Arg Ser Arg Phe Gln Val Leu
Val Ser Leu Val Phe Trp Asp Gln Cys Pro Gln Phe Val Gln Val Gln
Asp Trp Leu Pro Glu Met Ser Leu Leu Val Glu Val Ser His Thr
                 85
                                     90
Asn Leu Thr Glu Ile Thr Trp Met Val Phe Ile His Val Asn Ser Val
            100
                                105
                                                    110
```

<210> 99

<211> 1461

130

115

<212> DNA

<213> Saccharomyces cerevisiae

Val Met Leu Thr Thr Gly His Thr Ser Thr Thr Gly Val Leu Ser Val

125

140

120

Leu Thr Asp Thr Thr Phe Thr Gly Cys Thr Lys Lys Ile Leu

```
<400> 99
aacaagctat aatattgtta aatatagttg atcaacagca ttgtaatgat tacaagagac 60
gaggtggaat gaaccttatg aaatgcgtat tatatataaa ctgtaataag agctaagttg 120
aattgaaatc tacgatactt gatgttgaca ttatagcact agttcccagg aaaccctttc 180
gaaaaacaca gcaaaaacaa gagtactgta accaatgtaa catctgtaca ccagggaccc 240
acacattacc aaaatcaaaa ttatttttct aatgcctgtt atttttccta tttttcctct 300
ggcgcgtgaa tagcccgcag agacgcaaac aattttcctc gcagtttttc gcttgtttaa 360
tgcgtatttt cccagatagg ttcaaacctt tcatctgtat cccgtatatt taagatggcg 420
tttgctttct ccgttgattt ttttccttct tagtgatttt tttgcattaa atcccagaac 480
aatcatccaa ctaatcaaga atgccttcca gattcactaa gactagaaag cacagaggtc 540
acgtctcagg tatgtagttc catttggaag agggaatgaa agaaccaaga cggtgacttt 600
ttttttagtg ttgtgcaacc aatatgtcgt gtgtatatca tggtacagga gaatgtcaat 660
cagctaagtg tactcaacat atttctttgt gttttgattg cgaactttgt attaccatct 720
cactgttgag acggcttatt tgaggtaata gctcgagtaa atgtactctt ccatcgcaaa 780
ctgagcaaaa agaaagtgtg catagccttt gtcatacttc tcctttatta taccatgata 840
ttcagaacag tcatactgtc tactcatttt acggctataa aaggtaactt tcatttagat 900
tatggaaagc actaattatc gctgtatcaa atggttgtag agagcgcaat tatgaaaaag 960
agttaccacg tttcttttgt ttcgataaaa tgtccagttg aaaacctgtt ttactaacga 1020
tttaaaaatt gtatttcatt acaatatttt ttttgtacag ccggtaaagg tcgtatcggt 1080
aagcacagaa agcacccgg tggtagaggt atggccggtg gtcaacatca ccacagaatt 1140
aacatggata aataccatcc aggttatttc ggtaaggttg gtatgagata cttccacaag 1200
caacaagete atttetggaa gecagtettg aaettggaca aattgtggac attgateeca 1260
gaagacaaga gagaccaata cttgaaatct gcttctaagg aaactgctcc agttattgac 1320
actttggcag ccggttacgg taagatcttg ggtaagggta gaattccaaa tgttccagtt 1380
atcgtcaaag ctagattcgt ctccaagttg gctgaagaaa aaatcagagc tgctggtggt 1440
gttgttgaat tgatcgctta a
                                                                  1461
<210> 100
<211> 149
<212> PRT
<213> Saccharomyces cerevisiae
<400> 100
Met Pro Ser Arg Phe Thr Lys Thr Arg Lys His Arg Gly His Val Ser
 1
                  5
                                     10
                                                         15
Ala Gly Lys Gly Arg Ile Gly Lys His Arg Lys His Pro Gly Gly Arg
             20
                                 25
Gly Met Ala Gly Gly Gln His His Arg Ile Asn Met Asp Lys Tyr
                             40
```

His Pro Gly Tyr Phe Gly Lys Val Gly Met Arg Tyr Phe His Lys Gln

Gln Ala His Phe Trp Lys Pro Val Leu Asn Leu Asp Lys Leu Trp Thr

60

02-07-1999 EP99870141.1 SEQL

```
Leu Ile Pro Glu Asp Lys Arg Asp Gln Tyr Leu Lys Ser Ala Ser Lys

85

90

95
```

Glu Thr Ala Pro Val Ile Asp Thr Leu Ala Ala Gly Tyr Gly Lys Ile 100 105 110

Leu Gly Lys Gly Arg Ile Pro Asn Val Pro Val Ile Val Lys Ala Arg 115 120 125

Phe Val Ser Lys Leu Ala Glu Glu Lys Ile Arg Ala Ala Gly Gly Val 130 135 140

Val Glu Leu Ile Ala 145

<210> 101 <211> 1880 <212> DNA <213> Saccharomyces cerevisiae

<40.0> 101

```
ttttggtact tgaatttctc gtttctacta gctggattgc ttgtcctttt acaattcctg 60
ggagtttggc tattgctacc cttggtctta gcaccacttt tttccgctat ttgtatttgt 120
tgaactaatt tagtatctat ttccatttca ttataattca cgtttttagc agcctctctt 180
cttctaggta attggaaatc ttcttcttta ggcatagcgt gaccagctga aaggcagatg 240
tagtcctttt gaaagcaata tcataaaata tatatagttc tccatgatgt tcgggtcagt 360
cgctccgaag cgtaacctag tataataaat agttcattgc agaaaataac gaaagaaatg 420
gtggaatacg atctgttata tctaaactaa agctaactaa cggaataagc aaatacgaat 480
cgaccgctaa tttaacaaat atggttttag caatggaaag tagagtggca ccggaaattc 540
ctgggctcat tcaacctggg aatgtcacgc aagacttgaa gatgatggtc tgtaaattat 600
tgaattcccc aaaacctacg aaaacattcc ctggttccca gcctgtgtcc tttcagcatt 660
ctgatgtgga agagaagctg cttgcgcatg attactacgt ttgtgagaaa acagatggtc 720
tgcgggtgtt gatgtttata gtgataaatc ctgtgacggg tgagcaagga tgctttatga 780
ttgataggga aaataactat tatctggtta atggatttag gtttcccaga ttaccccaaa 840
agaagaaaga agagctgcta gagactcttc aagatggcac cttattagat ggtgaacttg 900
tcatacaaac taacccaatg acaaaattac aagagttgcg ttatttaatg ttcgattgtc 960
ttgctatcaa tggtagatgt ctcacacaat caccaacaag ttctagacta gcccaccttg 1020
gaaaagaatt ttttaaacca tacttegatt taagageage gtaceetaat egttgtaeta 1080
cttttccgtt caaaatttcc atgaaacata tggatttcag ttaccaatta gtaaaagttg 1140
ctaaaagttt agataaacta ccacatcttt ctgatggtct gatatttact cctgtgaagg 1200
caccttacac tgccggcgga aaagattcat tgttattaaa atggaagcca gaacaagaaa 1260
acaccgtgga cttcaaattg attttagata tcccaatggt ggaggatect tctttgccta 1320
aagatgatcg gaacaggtgg tattacaatt atgacgttaa gccagttttc agcttatatg 1380
```

Printed:17-08-2000 123

```
tctggcaagg cggagctgat gtcaattcac gtttaaaaca tttcgaccag cctttcgata 1440 ggaaggaatt tgaaatatta gaaagaacat acagaaaatt tgcagagttg agcgtttcag 1500 atgaggaatg gcaaaatttg aagaacctag aacagccatt aaatggtaga atagtagagt 1560 gcgcaaaaaa ccaagagact ggggcgtggg aaatgttaag attcagggat gataagttaa 1620 atggtaatca tacatcggtg gtccagaaag ttttggagag tatcaacgat tcagttcat 1680 tggaggacct cgaggaaatt gttggtgata ttaaaaggtg ctgggacgag agaagagcaa 1740 atatggctgg tggtagtgg agaccactac cgtctcaaag tcaaaatgcg acattatcta 1800 cctctaagcc agtccattca cagccccaa gtaatgataa ggagccaaaa tatgtagacg 1860 aggatgattg gtcggattag
```

<210> 102

<211> 459

<212> PRT

<213> Saccharomyces cerevisiae

<400> 102

Met Val Leu Ala Met Glu Ser Arg Val Ala Pro Glu Ile Pro Gly Leu
1 5 10 15

Ile Gln Pro Gly Asn Val Thr Gln Asp Leu Lys Met Met Val Cys Lys
20 25 30

Leu Leu Asn Ser Pro Lys Pro Thr Lys Thr Phe Pro Gly Ser Gln Pro 35 40 45

Val Ser Phe Gln His Ser Asp Val Glu Glu Lys Leu Leu Ala His Asp 50 55 60

Tyr Tyr Val Cys Glu Lys Thr Asp Gly Leu Arg Val Leu Met Phe Ile 65 70 75 80

Val Ile Asn Pro Val Thr Gly Glu Gln Gly Cys Phe Met Ile Asp Arg 85 90 95

Glu Asn Asn Tyr Tyr Leu Val Asn Gly Phe Arg Phe Pro Arg Leu Pro 100 105 110

Gln Lys Lys Glu Glu Leu Leu Glu Thr Leu Gln Asp Gly Thr Leu 115 120 125

Leu Asp Gly Glu Leu Val Ile Gln Thr Asn Pro Met Thr Lys Leu Gln 130 135 140

Glu Leu Arg Tyr Leu Met Phe Asp Cys Leu Ala Ile Asn Gly Arg Cys 145 150 155 160

Leu Thr Gln Ser Pro Thr Ser Ser Arg Leu Ala His Leu Gly Lys Glu



- Phe Phe Lys Pro Tyr Phe Asp Leu Arg Ala Ala Tyr Pro Asn Arg Cys 180 185 190
- Thr Thr Phe Pro Phe Lys Ile Ser Met Lys His Met Asp Phe Ser Tyr 195 200 205
- Gln Leu Val Lys Val Ala Lys Ser Leu Asp Lys Leu Pro His Leu Ser 210 215 220
- Asp Gly Leu Ile Phe Thr Pro Val Lys Ala Pro Tyr Thr Ala Gly Gly 225 230 235 240
- Lys Asp Ser Leu Leu Leu Lys Trp Lys Pro Glu Gln Glu Asn Thr Val 245 250 255
- Asp Phe Lys Leu Ile Leu Asp Ile Pro Met Val Glu Asp Pro Ser Leu 260 265 270
- Pro Lys Asp Asp Arg Asn Arg Trp Tyr Tyr Asn Tyr Asp Val Lys Pro 275 . 280 285
- Val Phe Ser Leu Tyr Val Trp Gln Gly Gly Ala Asp Val Asn Ser Arg 290 295 300
- Leu Lys His Phe Asp Gln Pro Phe Asp Arg Lys Glu Phe Glu Ile Leu 305 310 315 320
- Glu Arg Thr Tyr Arg Lys Phe Ala Glu Leu Ser Val Ser Asp Glu Glu 325 330 335
- Trp Gln Asn Leu Lys Asn Leu Glu Gln Pro Leu Asn Gly Arg Ile Val
- Glu Cys Ala Lys Asn Gln Glu Thr Gly Ala Trp Glu Met Leu Arg Phe 355 360 365
- Arg Asp Asp Lys Leu Asn Gly Asn His Thr Ser Val Val Gln Lys Val 370 375 380
- Leu Glu Ser Ile Asn Asp Ser Val Ser Leu Glu Asp Leu Glu Glu Ile 385 390 395 400
- Val Gly Asp Ile Lys Arg Cys Trp Asp Glu Arg Arg Ala Asn Met Ala 405 410 415
- Gly Gly Ser Gly Arg Pro Leu Pro Ser Gln Ser Gln Asn Ala Thr Leu

425

430

Ser Thr Ser Lys Pro Val His Ser Gln Pro Pro Ser Asn Asp Lys Glu
435 440 445

Pro Lys Tyr Val Asp Glu Asp Asp Trp Ser Asp 450 455

<210> 103

<211> 1076

<212> DNA

<213> Saccharomyces cerevisiae

<400> 103

caatattata gttactacct atttgattat taagatcaaa agtaatcctt catctcatat 60 caaaaagaaa ggtttgtaaa agagcattgg gcagagagta tttgtcctgc gtatagagga 120 gaaaaaattg tactgatgct aattttggtg tcgttcttct tttttatttt gtaagactgt 180 tttccagaaa tgtttgggtt tatttttaa ttttttgaaa cattttttc atcctttctc 240 attttgtcat ttcatttttc tgtggaaaat ttcactgacg cgaagaagcg atgaaatttc 300 caacatcctc ccatcatccc aatattggca tacacacaca tgcagcacag cggaactgcg 360 gaggtcagag gcaatgtggc agagacgctg gcgcgcctgt attgtataat agtatatttt 420 acactcaatt caattttttg atattaaatt agtgtgtaaa aagcttctga aatcaagaag 480 cccgtaccag aagttcaatc atgaaataca tccaaactga acaacaaatc gaagtcccag 540 aaggtgtcac tgtcagcatc aagtccagaa tcgtcaaggt tgttggtcca agaggtactt 600 tgaccaagaa cttgaagcac attgatgtta ccttcaccaa ggtcaacaac caattgatca 660 aggttgctgt tcacaacggt ggcagaaagc acgttgctgc tttgagaacc gtcaagtctt 720 tggttgacaa catgatcact ggtgtcacca agggttacaa gtacaagatg agatacgtct 780 acgcgcattt cccaatcaac gtcaacattg ttgaaaagga tggtgccaaa ttcattgaag 840 tcagaaactt tttgggtgac aagaagatca gaaacgttcc agttagagat ggtgttacta 900 tttcccaaaa cgctgctgac ttgcaacaaa tctgtcgtgt cagaaacaag gatatccgta 1020 aatttttgga cggtatctac gtttctcaca agggttttat tactgaagat ttataa 1076

<210> 104

<211> 191

<212> PRT

<213> Saccharomyces cerevisiae

<400> 104

Met Lys Tyr Ile Gln Thr Glu Gln Gln Ile Glu Val Pro Glu Gly Val
1 5 10 15

Thr Val Ser Ile Lys Ser Arg Ile Val Lys Val Val Gly Pro Arg Gly 20 25 30



Thr Leu Thr Lys Asn Leu Lys His Ile Asp Val Thr Phe Thr Lys Val 35 40 45

Asn Asn Gln Leu Ile Lys Val Ala Val His Asn Gly Gly Arg Lys His 50 55 60

Val Ala Ala Leu Arg Thr Val Lys Ser Leu Val Asp Asn Met Ile Thr 65 70 75 80

Gly Val Thr Lys Gly Tyr Lys Tyr Lys Met Arg Tyr Val Tyr Ala His
85 90 95

Phe Pro Ile Asn Val Asn Ile Val Glu Lys Asp Gly Ala Lys Phe Ile 100 105 110

Glu Val Arg Asn Phe Leu Gly Asp Lys Lys Ile Arg Asn Val Pro Val 115 120 125

Arg Asp Gly Val Thr Ile Glu Phe Ser Thr Asn Val Lys Asp Glu Ile 130 135 140

Val Leu Ser Gly Asn Ser Val Glu Asp Val Ser Gln Asn Ala Ala Asp 145 150 155 160

Leu Gln Gln Ile Cys Arg Val Arg Asn Lys Asp Ile Arg Lys Phe Leu 165 170 175

Asp Gly Ile Tyr Val Ser His Lys Gly Phe Ile Thr Glu Asp Leu 180 185 190

<210> 105

<211> 1694

<212> DNA

<213> Saccharomyces cerevisiae

<400> 105

aatcaaaacg atcgttatac aattcattga ggaacctctc tagtccgatc cgtctttcat 60 cgatcattc cggatcatca tatcttctt gccacctcct gtccaataca ccgggctttt 120 cagggaagtc atatgggatg gtgcttccta catctctct caatcgtgtc ttcagtttcc 180 aaaactcgga ataccttttg taaaggcgct tgtttggtgt actaacaccg tataaaacat 240 acttggggtt gatttaaca tcgtccacct tgattcttaa cttttcactc attttcccta 300 cagaattagc tgccatcaat atcaattatc aacccttata tgactttatt gttttgttt 360 ttgttggcta attaattgat acaaatcttt aggcgaaaaa taaaaaaaa ataaagtaaa 420 gaaggaaaaa ttaggcgata ttaaaacaaa tctaaaataa agacaagaaa cgaaaaagag 480 gttaatcaag tgatatttc tcggtttctg cttgcaattc atttaccggta agttgtccag 600

```
gtgacggtta cttaaaggtg tgggataata agctgttaga taatgaaaat ccaaaagata 660
agtcatatte teaetttgte cataagteeg gattgeacea tgtegatgte ttgeaageta 720
ttgagagaga tgcatttgaa ttatgccttg ttgctaccac ttcattttct ggcgatttac 780
tcttctatcg tatcactaga gaagatgaga ctaaaaaaagt tatattcgag aaattggatc 840
ttctagactc agacatgaaa aagcattcct tttgggcatt aaaatggggt gcctcaaatg 900
acagactact ttcccatagg ctggttgcta cagacgtcaa agggaccact tacatttgga 960
agtttcaccc gtttgcagat gagtcaaatt ctttaacact aaattggagc cccacgttag 1020
aattacaagg cactgtcgaa tcgcctatga ctccaagtca atttgccact tctgtggata 1080
tctctgaacg aggactaatt gccacaggtt ttaataatgg aacagtacaa atttcagaac 1140
tatctacatt acgcccgttg tacaattttg aatctcagca ttctatgatt aataattcga 1200
attecateag ateggtgaaa tttteteete aaggateett attageeatt geteaegatt 1260
caaattcatt tggttgcatc actctatatg aaactgaatt tggtgaaaga ataggctcct 1320
tatccgtacc aacccatage tegcaggeaa gtetgggtga atttgcacat tetagetggg 1380
tcatgagtct atcgtttaat gattctggtg aaacattatg cagtgccgga tgggatggta 1440
aattgagatt ttgggatgta aaaacaaagg aaagaatcac tacattgaat atgcattgtg 1500
atgatattga aattgaagag gatatcttag ctgttgatga acatggagat tctttagctg 1560
aacctggtgt ctttgacgtg aagtttttga aaaaaggttg gagatctggt atgggagctg 1620
atttaaatga aagtttatgc tgtgtttgtt tagatagaag catcaggtgg tttagagaag 1680
ctggcggtaa ataa
                                                                  1694
```

```
<210> 106
```

<211> 397

<212> PRT

<213> Saccharomyces cerevisiae

<400> 106

Met Ser Lys Val Phe Ile Ala Thr Ala Asn Ala Gly Lys Ala His Asp 1 5 10 15

Ala Asp Ile Phe Ser Val Ser Ala Cys Asn Ser Phe Thr Val Ser Cys
20 25 30

Ser Gly Asp Gly Tyr Leu Lys Val Trp Asp Asn Lys Leu Leu Asp Asn 35 40 45

Glu Asn Pro Lys Asp Lys Ser Tyr Ser His Phe Val His Lys Ser Gly
50 55 60

Leu His His Val Asp Val Leu Gln Ala Ile Glu Arg Asp Ala Phe Glu 65 70 75 80

Leu Cys Leu Val Ala Thr Thr Ser Phe Ser Gly Asp Leu Leu Phe Tyr 85 90 95

Arg Ile Thr Arg Glu Asp Glu Thr Lys Lys Val Ile Phe Glu Lys Leu 100 105 110





- Asp Leu Leu Asp Ser Asp Met Lys Lys His Ser Phe Trp Ala Leu Lys 115 120 125
- Trp Gly Ala Ser Asn Asp Arg Leu Leu Ser His Arg Leu Val Ala Thr 130 135 140
- Asp Val Lys Gly Thr Thr Tyr Ile Trp Lys Phe His Pro Phe Ala Asp 145 150 155 160
- Glu Ser Asn Ser Leu Thr Leu Asn Trp Ser Pro Thr Leu Glu Leu Gln 165 170 175
- Gly Thr Val Glu Ser Pro Met Thr Pro Ser Gln Phe Ala Thr Ser Val 180 185 190
- Asp Ile Ser Glu Arg Gly Leu Ile Ala Thr Gly Phe Asn Asn Gly Thr 195 200 205
- Val Gln Ile Ser Glu Leu Ser Thr Leu Arg Pro Leu Tyr Asn Phe Glu 210 215 220
- Ser Gln His Ser Met Ile Asn Asn Ser Asn Ser Ile Arg Ser Val Lys 225 230 235 240
- Phe Ser Pro Gln Gly Ser Leu Leu Ala Ile Ala His Asp Ser Asn Ser 245 250 255
- Phe Gly Cys Ile Thr Leu Tyr Glu Thr Glu Phe Gly Glu Arg Ile Gly 260 265 270
- Ser Leu Ser Val Pro Thr His Ser Ser Gln Ala Ser Leu Gly Glu Phe 275 280 285
- Ala His Ser Ser Trp Val Met Ser Leu Ser Phe Asn Asp Ser Gly Glu 290 295 300
- Thr Leu Cys Ser Ala Gly Trp Asp Gly Lys Leu Arg Phe Trp Asp Val 305 310 315 320
- Lys Thr Lys Glu Arg Ile Thr Thr Leu Asn Met His Cys Asp Asp Ile 325 330 335
- Glu Ile Glu Glu Asp Ile Leu Ala Val Asp Glu His Gly Asp Ser Leu 340 345 350
- Ala Glu Pro Gly Val Phe Asp Val Lys Phe Leu Lys Lys Gly Trp Arg 355 360 365

```
Ser Gly Met Gly Ala Asp Leu Asn Glu Ser Leu Cys Cys Val Cys Leu 370 375 380
```

```
Asp Arg Ser Ile Arg Trp Phe Arg Glu Ala Gly Gly Lys
385 390 395
```

```
<210> 107
```

<211> 1037

<212> DNA

<213> Saccharomyces cerevisiae

```
<400> 107
```

```
ctggttccaa ccatatctta tgggaagatc tatcaggaaa ccttagaatc ttagcttcga 60
tggagggaca atagcggggg cctttgatgg tcgtgtcctg aatatgaatg gattgatgca 120
agttattgcg caaaaagtcg tgcatttgag gtgtggtatg tgtgccaaag caatccagtt 180
gcttcgtggg ttcgactgac acggtttcat tcagaaaact catagggaca ggcaacgcat 240
ctcccttctg gacctccaga gcgctgaaat caatggattc cttggccagc ctagccggag 300
tgcctgtttt caaacgccct aactgaaagc ccacctcatt ttgtagagta ttgctgatcc 360
catatgttgg ctgctcgcca attcttcctg ctgcaatacg cttgtcgccg atgtgaattt 420
ctgcactgag aaacgtacct gtagtgatta tgacctgatc tgccccaacc tgggtaccgt 480
catccagaac cacacetttg atgacettgt ggccacatec ggggtcatac aagatcaagt 540
cagcaacttt gttctgcagc agagacaagt tggggtgtgc tttcttgtcg gaaagttccc 600
tttgcatgta tttcttatat aactctctgt ctatctgggc tctgggcccc cacacagcag 660
gaccettget tetgttcage attttgaatt geactecage gagateagtt actttgccca 720
tcagcccatc gagagcgtcg atttccttta caaggatgcc cttacccact ccaccaatag 780
aggggttgca cgaacactta ccaatatctg ttaacgatgg tgtaatgaga gtagtatgcg 840
caccagtect agatgacgca getgeagett egeageetge atgtecagea eegatgacca 900
cgacctgtgt cttggttgtg ggttggaagc ttgtcaaaga agatatggtc aaccgcctcc 960
tgaggacctg taagggaaat gaagtacacg aagacgccaa ggttgttaca cgcagcatcg 1020
ttctttgggg cgtttag
                                                                  1037
```

```
<210> 108
```

<211> 178

<212> PRT

<213> Saccharomyces cerevisiae

## <400> 108

Met Thr Leu Trp Pro His Pro Gly Ser Tyr Lys Ile Lys Ser Ala Thr
1 5 10 15

Leu Phe Cys Ser Arg Asp Lys Leu Gly Cys Ala Phe Leu Ser Glu Ser 20 25 30

Ser Leu Cys Met Tyr Phe Leu Tyr Asn Ser Leu Ser Ile Trp Ala Leu 35 40 45





Gly Pro His Thr Ala Gly Pro Leu Leu Leu Phe Ser Ile Leu Asn Cys
50 55 60

Thr Pro Ala Arg Ser Val Thr Leu Pro Ile Ser Pro Ser Arg Ala Ser 65 70 75 80

Ile Ser Phe Thr Arg Met Pro Leu Pro Thr Pro Pro Ile Glu Gly Leu 85 90 95

His Glu His Leu Pro Ile Ser Val Asn Asp Gly Val Met Arg Val Val
100 105 110

Cys Ala Pro Val Leu Asp Asp Ala Ala Ala Ala Ser Gln Pro Ala Cys 115 120 125

Pro Ala Pro Met Thr Thr Cys Val Leu Val Val Gly Trp Lys Leu
130 135 140

Val Lys Glu Asp Met Val Asn Arg Leu Leu Arg Thr Cys Lys Gly Asn 145 150 155 160

Glu Val His Glu Asp Ala Lys Val Val Thr Arg Ser Ile Val Leu Trp 165 170 175

Gly Val

<210> 109

<211> 731

<212> DNA

<213> Saccharomyces cerevisiae

# <400> 109

taccatggaa caccggtgat cattctggtc acttggtctg gggcaatacc agtcaacatg 60 gtggtgaagt caccgtagtt gaaaacggct tcagcaactt caactgggta ggtttccgtt 120 gggtgggcgg cttggaacat gtagtattga gccaagtgag ctctgatatc agagacgtag 180 acacctaatt caaccaaatt gactctttcg tcagactgag ctagagtggt ggttgcggaa 240 gcagtagcag cgatggcagc gacaccagcg gcgattgaag ttaatttgac cattgtattt 300 gttttgttt ttagtgctgg tataagctta acaggaaaga aagaaataaa gatatattct 360 caaaagcata cagttgaagc agctctattt atacccgttc ctctatcagt catcactact 420 taaacgattc gttaacagat gctcatttag cacctcacat atcctccata tctcatcttt 480 cacacaatct cattatcact atggagatgc tcttgtttct gaacgaatca tacatctttc 540 atagacttcg tatgtggga attgtattat ggcactcatg tgtattcgta tgcgcagaat 600 gtgggaatgc caattatagg gtgccgaggt gccttataaa acccttttct gtgcctgtga 660 catttccttt ttcggtcaaa aagaatatcc gaattttaga tttggaccct cgtacagaag 720

```
cttattgtta a
                                                                   731
<210> 110
<211> 76
<212> PRT
<213> Saccharomyces cerevisiae
<400> 110
Met Glu Met Leu Leu Phe Leu Asn Glu Ser Tyr Ile Phe His Arg Leu
                                      10
Arg Met Trp Ser Ile Val Leu Trp His Ser Cys Val Phe Val Cys Ala
                                 25
Glu Cys Gly Asn Ala Asn Tyr Arg Val Pro Arg Cys Leu Ile Lys Pro
                             40
Phe Ser Val Pro Val Thr Phe Pro Phe Ser Val Lys Lys Asn Ile Arg
                         55
Ile Leu Asp Leu Asp Pro Arg Thr Glu Ala Tyr Cys
                     70
<210> 111
<211> 1025
<212> DNA
<213> Saccharomyces cerevisiae
<400> 111
tecttaettt agtetattat caatatetet tececeteet aaatatgtae tettttattt 60
tttttaattg tgaaggaaca attcaagtta gaactctttt gataggaaac attatttcct 120
gtgtagccta atgtttaatg cctaattttt ttctaaaatg cagcaacata catatgttga 180
gtcgtataga catctatata taacaagcac agaaccgtct aattggtatt tttcaggaca 240
ttttaaacat ccgtacaacg agaacccata cattactttt tttaatattc tttttgtttt 300
caccgccttc tttttatttt tatccgaaga tcttttggaa cccgctctgc gaatagcgaa 360
gctaggatac caaattgaaa cttggacata actcatcatt aaagaagtat actgttaaga 420
gaggcattca tttcgtgtat tataacgttt agcatcagtt acccttgaaa gcccaacata 480
tacaaaaata cgcgtccaag atgtctacta aagcccaaaa ccctatgcgt gatttgaaga 540
tcgagaaatt ggtcttgaac atctccgttg gtgaatctgg tgacagatta accagagcct 600
ccaaggtttt agaacaatta tctggtcaaa ctccagttca atccaaggcc agatacactg 660
```

tcagaacttt cggtatcaga agaaacgaaa aaattgctgt tcacgttacc gtcagaggtc 720 caaaggctga agaaattttg gaaagaggtt tgaaggtcaa ggaataccaa ttgagagaca 780 gaaacttctc tgctaccggt aacttcggtt tcggtattga cgaacacatt gacttgggta 840 tcaagtatga cccatccatc ggtattttcg gtatggattt ctatgtcgtc atgaacagac 900 caggtgctag agtcactaga agaaagagat gtaagggtac tgttggtaac tcccacaaga 960

caactaagga agacaccgtc tcttggttca agcaaaagta cgacgctgat gtgctcgata 1020 aataa 1025

<210> 112

<211> 174

<212> PRT

<213> Saccharomyces cerevisiae

<400> 112

Met Ser Thr Lys Ala Gln Asn Pro Met Arg Asp Leu Lys Ile Glu Lys

1 5 10 15

Leu Val Leu Asn Ile Ser Val Gly Glu Ser Gly Asp Arg Leu Thr Arg
20 25 30

Ala Ser Lys Val Leu Glu Gln Leu Ser Gly Gln Thr Pro Val Gln Ser 35 40 45

Lys Ala Arg Tyr Thr Val Arg Thr Phe Gly Ile Arg Arg Asn Glu Lys
50 55 60

Ile Ala Val His Val Thr Val Arg Gly Pro Lys Ala Glu Glu Ile Leu 65 70 75 80

Glu Arg Gly Leu Lys Val Lys Glu Tyr Gln Leu Arg Asp Arg Asn Phe 85 90 95

Ser Ala Thr Gly Asn Phe Gly Phe Gly Ile Asp Glu His Ile Asp Leu 100 105 110

Gly Ile Lys Tyr Asp Pro Ser Ile Gly Ile Phe Gly Met Asp Phe Tyr 115 120 125

Val Val Met Asn Arg Pro Gly Ala Arg Val Thr Arg Arg Lys Arg Cys 130 135 140

Lys Gly Thr Val Gly Asn Ser His Lys Thr Thr Lys Glu Asp Thr Val 145 150 155 160

Ser Trp Phe Lys Gln Lys Tyr Asp Ala Asp Val Leu Asp Lys 165 170

<210> 113

<211> 1258

<212> DNA

```
<400> 113
agccatgtcg gtcgcactag acttttcttc tcactgtcac ttacctgttt gaaatcatgt 60
ccttttttt tttgccattc ttatacattt tctttccttc tgaaattaac tgtacaccca 120
taccctatat acacccatac cctattttta aatataaaaa gtaaacttca ttttgaaaga 180
ccactctgca tcagcacgcg ggctctggaa ggaagaaatg acgtttcggc ggaataccct 240
ttcagaaggt ctgctcttgt ggctggttca tgggagacac ccagcggagc tcctcccgag 300
aaaggcccct tcatctctgc cgattgctga cggaaagcag tagcggaggt ttgagttctc 360
tacgccgaga gtacactgcc gtaatatcac aatgtttcga ctaacggtta cagtacgtta 420
aattagatac tgcctatgaa ttgacatatt agataatgtc aaattttaca aaaacctaag 480
acaacaggaa ataaacaaag atgggtaaag gtaagccaag aggtttgaac tctgctagaa 540
agctacgtgt ccacagaaga aacaagtatg ttgactattt caaaattaaa aaaaactatc 600
aaccccctat tgtgatatcg ttttaggtga aggaaatgtt gtgagctctg gagtgataaa 660
tttatcaagt aacatatcct ggcgcaaatc agtttggaga ggcttaaaat gacacgtcac 720
agtgataaaa agtaatgaat agtgaacggt cagcttcggc cattcttccc aatctatagt 780
gtggaaaata aaccttttct tcccaaaata actcagaaag tcacaggagg ccgtttttta 840
caacggaatc attitttac taacagtitt titttattat tatagccgtt gggccgaaaa 900
```

caactacaag aagagattgt tgggtactgc cttcaagtct tctccattcg gtggttcttc 960 tcatgccaag ggtatcgtct tggaaaaatt gggtatcgaa tccaagcaac ctaactctgc 1020 tatcagaaag tgtgttagag ttcaattaat caagaacggt aagaaggtca ctgctttcgt 1080 tccaaacgat ggttgtttga actttgtcga cgaaaatgat gaagtcttgc tagcaggttt 1140 cggtagaaag ggtaaagcta agggtgatat tccaggtgtt agattcaagg tcgttaaggt 1200

<210> 114 <211> 145 <212> PRT <213> Saccharomyces cerevisiae

<213> Saccharomyces cerevisiae

<400> 114

Met Gly Lys Gly Lys Pro Arg Gly Leu Asn Ser Ala Arg Lys Leu Arg 1 5 10 15

ctctggtgtc tccttgttgg ctttgtggaa agaaaagaag gaaaagccaa gatcataa

Val His Arg Arg Asn Asn Arg Trp Ala Glu Asn Asn Tyr Lys Lys Arg
20 25 30

Leu Leu Gly Thr Ala Phe Lys Ser Ser Pro Phe Gly Gly Ser Ser His
35 40 45

Ala Lys Gly Ile Val Leu Glu Lys Leu Gly Ile Glu Ser Lys Gln Pro 50 55 60

Asn Ser Ala Ile Arg Lys Cys Val Arg Val Gln Leu Ile Lys Asn Gly 65 70 75 80

Lys Lys Val Thr Ala Phe Val Pro Asn Asp Gly Cys Leu Asn Phe Val

02-07-1999

EP99870141.1

SEQL

85 90 95

Asp Glu Asn Asp Glu Val Leu Leu Ala Gly Phe Gly Arg Lys Gly Lys
100 105 110

Ala Lys Gly Asp Ile Pro Gly Val Arg Phe Lys Val Val Lys Val Ser 115 120 125

Gly Val Ser Leu Leu Ala Leu Trp Lys Glu Lys Lys Glu Lys Pro Arg 130 135 140

Ser 145

<210> 115 <211> 1733

<212> DNA

<213> Saccharomyces cerevisiae

<400> 115

ggactacttt acagggtaat gaatatttgg gcgtttttcg ctattttagc atgctgtagt 60 gtatgtactg tgcatcgtca tgtagcacta tttcagccgt atttttcttt ttttctttcg 120 caccgtctgt ggttgtaaag ttactgacac ttttttttct agaaagttcc ggaaaattgc 180 gacacteggt ggagetegag agttgtatee agttttettg tteggegata tteegaacea 240 ggtcgggttg ggctaacagc cgcccaggat ggaagaatta agaatttcat agaagccttc 300 agttettgge gaagtaaagt ggeaaaacaa atggaagate tattgeatta catatataaa 360 agcattagaa caatcttttc tcattgacag gtattctcat tgctctatat atattttctt 420 cttcttgaaa gaaatatcag tattacaatc ataacaacaa ccaaaagaaa ataactaata 480 gaccccatta caatatagaa atgttttcca tattcaattc accatgtgtt tttgaacagc 540 tgccatcttt tagtcagccc ctacattcgc gttattttga ttgcagttct ccagtgagct 600 attatccaga atgtaaaagg aggaaagcaa taaaagctaa cctaagagct ccaaaaaaaa 660 gcgatgcaaa ttgttcagaa cctttgaggt atgcacttgc tgaaacacca aatggttata 720 cattaagctt gtctaagcgg attccatatg aacttttttc aaagtacgtt aatgagaaat 780 taggtgagct aaaggagaac cattacagac caacttacca tgttgtccaa gatttttttg 840 gaaaccagta ttatgttgaa gatgaagcgg atgaagatgc tctattgaga tctgcattga 900 aagatctgga ttttagagcc ataggaaaga aaattgctaa ggatcttttc caagactacg 960 aaatagaatt gaatcataga ggtgatgaat tgagcatatt gagtaagaag gataaaatct 1020 ttaaggaatt ctctctagac caagtgtttg aagatgtttt tgttattggc tgtggagttg 1080 aaaacataga tgatggctcg agagaaaaat atgcactttt aaagattggt ttagttaagc 1140 atgaggaaga aatttccgaa ggtggcatca acgaaccaaa gatgccaata attgaatcca 1200 aaatagacga gtctcacgat gatgttaaca tgtctgaatc tttgaaggag gaagaagcgg 1260 agaaagcgaa agaaccacta accaaagaag accaaataaa aaaatggata gaggaagaaa 1320 gattgatgca ggaggaaagc agaaaatcag aacaggaaaa agctgccaag gaagatgaag 1380 aaaggcaaaa gaaagagaag gaagccagat tgaaggcaag gaaagaatct ttgataaata 1440 agcaaaaaac caagaggtcc cagcaaaaaa aattgcaaaa ttccaaatca ttgcctatct 1500 ctgagattga ggccagcaat aaaaataata atagcaattc tggttcagca gaaagtgata 1560

135

Printed:17-08-2000

atgaaagtat aaacagtgat tétgatacga etttggattt etetgtgtet ggtaatacae 1620 taaaaaaaca cgcttcaccc ctattagaag acgttgagga tgaggaagtt gacagataca 1680 acgagtccct aagcagatct cccaagggaa actctattat tgaggagata taa <210> 116 <211> 410 <212> PRT <213> Saccharomyces cerevisiae <400> 116 Met Phe Ser Ile Phe Asn Ser Pro Cys Val Phe Glu Gln Leu Pro Ser 10 Phe Ser Gln Pro Leu His Ser Arg Tyr Phe Asp Cys Ser Ser Pro Val 20 25 Ser Tyr Tyr Pro Glu Cys Lys Arg Lys Ala Ile Lys Ala Asn Leu 35 40 . Arg Ala Pro Lys Lys Ser Asp Ala Asn Cys Ser Glu Pro Leu Arg Tyr 55 Ala Leu Ala Glu Thr Pro Asn Gly Tyr Thr Leu Ser Leu Ser Lys Arg 65 70 75 Ile Pro Tyr Glu Leu Phe Ser Lys Tyr Val Asn Glu Lys Leu Gly Glu 90 Leu Lys Glu Asn His Tyr Arg Pro Thr Tyr His Val Val Gln Asp Phe

Leu Lys Giu Ash His Tyr Arg Pro Thr Tyr His Vai Vai Gin Asp Phe 100 105 110

Phe Gly Asn Gln Tyr Tyr Val Glu Asp Glu Ala Asp Glu Asp Ala Leu 115 120 125

Leu Arg Ser Ala Leu Lys Asp Leu Asp Phe Arg Ala Ile Gly Lys Lys
130 . 135 140

Gly Asp Glu Leu Ser Ile Leu Ser Lys Lys Asp Lys Ile Phe Lys Glu 165 170 175

Phe Ser Leu Asp Gln Val Phe Glu Asp Val Phe Val Ile Gly Cys Gly 180 185 190

Val Glu Asn Ile Asp Asp Gly Ser Arg Glu Lys Tyr Ala Leu Leu Lys

200 -

205

Ile Gly Leu Val Lys His Glu Glu Glu Ile Ser Glu Gly Gly Ile Asn 210 215 220

Glu Pro Lys Met Pro Ile Ile Glu Ser Lys Ile Asp Glu Ser His Asp 225 230 235 240

Asp Val Asn Met Ser Glu Ser Leu Lys Glu Glu Glu Ala Glu Lys Ala 245 250 255

Lys Glu Pro Leu Thr Lys Glu Asp Gln Ile Lys Lys Trp Ile Glu Glu 260 265 270

Glu Arg Leu Met Gln Glu Glu Ser Arg Lys Ser Glu Gln Glu Lys Ala 275 280 285

Ala Lys Glu Asp Glu Glu Arg Gln Lys Lys Glu Lys Glu Ala Arg Leu 290 295 300

Lys Ala Arg Lys Glu Ser Leu Ile Asn Lys Gln Lys Thr Lys Arg Ser 305 310 315 320

Gln Gln Lys Lys Leu Gln Asn Ser Lys Ser Leu Pro Ile Ser Glu Ile 325 330 335

Glu Ala Ser Asn Lys Asn Asn Asn Ser Asn Ser Gly Ser Ala Glu Ser 340 345 350

Asp Asn Glu Ser Ile Asn Ser Asp Ser Asp Thr Thr Leu Asp Phe Ser 355 360 365

Val Ser Gly Asn Thr Leu Lys Lys His Ala Ser Pro Leu Leu Glu Asp 370 375 380

Val Glu Asp Glu Glu Val Asp Arg Tyr Asn Glu Ser Leu Ser Arg Ser 385 390 395 400

Pro Lys Gly Asn Ser Ile Ile Glu Glu Ile 405 410

<210> 117

<211> 890

<212> DNA

<213> Saccharomyces cerevisiae

```
<400> 117
caaaaaagtt ttcggatgaa ccggattaat acaagtaaaa tcagcaaaga tatagaagac 60
aaaataagcg tgaaaacaat cataaaccac tcacaacggg ggttttcagc tgttactcct 120
ccatacatac attttgataa agatataatg ttatatttct tttcgtaatt ttgttttact 180
teggtttgct ctatagattt catcageege acegaaaagg gagateaata aggtaceett 240
taaaagggat aagaagccta catcacccca ataaatggag taatggccag cattggatga 300
agagaagaat tacgggatac tgggataaca ctgttaaaaa atgcttcgcg acgtgagggt 360
cttcttcata taaattgaac tgccaaatct ctttcacatt atccaggata gtttggaatg 420
tgtgttactg aaggatcaga atcaataaat acaatcaata caaatattta gcgcataaaa 480
ttcaaacaaa gtttactgaa atgaagttag attcaggaat atactcagag gcacaaagag 540
ttgtgagaac tccaaagttt agatatatta tgttagggct ggtgggcgct gctgtggtac 600
cgaccgcata catgaggaga ggctatacgg ttcctgcaca tagcttagac aacatcaacg 660
gcgtagacac aactaaggcg tctgttatgg gtacagaaca gagagcagct atgacgaagg 720
gtaagagttt acaagagatg atggatgatg atgaagtaac gtatttgatg ttcctcttca 780
atcatgtaag ggaatttgta cttggttccc tgcatttatg ttctttgcat tttgttttcg 840
catttaatca tagtacgaca aacggggaag gggattgtga ttttacataa
<210> 118
<211> 129
<212> PRT
<213> Saccharomyces cerevisiae
<400> 118
Met Lys Leu Asp Ser Gly Ile Tyr Ser Glu Ala Gln Arg Val Val Arg
                  5
                                                         15
Thr Pro Lys Phe Arg Tyr Ile Met Leu Gly Leu Val Gly Ala Ala Val
             20
                                 25
                                                     30
Val Pro Thr Ala Tyr Met Arg Arg Gly Tyr Thr Val Pro Ala His Ser
         35
                             40
Leu Asp Asn Ile Asn Gly Val Asp Thr Thr Lys Ala Ser Val Met Gly
                         55
                                             60
Thr Glu Gln Arg Ala Ala Met Thr Lys Gly Lys Ser Leu Gln Glu Met
                     70
                                         75
Met Asp Asp Glu Val Thr Tyr Leu Met Phe Leu Phe Asn His Val
                 85
                                     90
```

Arg Glu Phe Val Leu Gly Ser Leu His Leu Cys Ser Leu His Phe Val

Phe Ala Phe Asn His Ser Thr Thr Asn Gly Glu Gly Asp Cys Asp Phe

105

110

100

. 115

Thr

```
<210> 119
<211> 1418
<212> DNA
<213> Saccharomyces cerevisiae
<400> 119
aatctttgcg attgaattgc tgcacgaaca ttaacattag ttcttttgct aaagtttctc 60
caatatctgg aatatcagaa tttagcaagg caatcaaagc ggataaagca gggattagcc 120
tgccattctg tttgttcaga gtaaaatcca caacattttt acaaattatg ttgcggccaa 180
taagtatatt gacttgaaat aagtctctgt gcgattcctg taagttgtcc attgttaaat 240
tggatattat gggtgaaacg tgtgacctta tcatttccca gttttctctc tgaaatttaa 300
tgtcttcatc ctgtatggta gcggtagaca tgctggtttg tagccttttt attccttttt 360
aggtttctta ctcacaatgc caaaataaat atcagtgtaa tataattttt caagagtacg 420
taatggaaaa agataaaaat aaggaccgtc ataaaaagag acgtgattaa acctaaaaat 480
ctaaagtaaa gaagtgtaag atggttgagg aaaattccag agttttgatt gttcttcctt 540
atacaccgcc tagtgctact ttgcagagga ttatagggca aactattccg ttcttaagag 600
aatgtcaaag tcaactagac atcgtgattg tacctgaatt caaaacctca ttccagttgg 660
attctgcgct agggaagatg tacagtatta ccagggatgt ccttttgggc tatggaatga 720
tcaacagcgg aatcaacatc atattcaaca atattcattt cgtcgagagt aatttgcaat 780
ggaaagtggt tttattgcca caggaatcca cttttgaaac ttggaagcta gagttgggac 840
aaggacaata ccatagtata gaacattatg cattacacga taatataatg gaagagatag 900
aaggtcccaa agatgctaac aaatttcatg tcaccgcatt gggcggaacg ttcgaccaca 960
ttcacgatgg acataaaata ttgttgagcg tctctacatt catcacgtca caaaggttaa 1020
tttgtggaat tacgtgcgat gagctcttgc aaaacaagaa atacaaagag ttgattgaac 1080
cttatgatac acgatgcagg cacgtacatc aattcatcaa gttgttaaaa ccggatctct 1140
ccgtagaact agttccctta agggacgtgt gcggcccac agggaaagta cccgagatag 1200
aatgtttagt tgtgagtaga gaaaccgtca gtggggcaga gactgtgaat aagactagga 1260
ttgaaaaagg catgagcca ttggcagtac atgtggttaa tgtacttgga ggaagggagg 1320
aaqacggctg gagcgagaag ttaagcagca cggaaatcag acgcctactt aagtcctctg 1380
                                                                  1418
cttcgccaac gtgcactcca caaaaccctt gcgtataa
<210> 120
<211> 305
<212> PRT
<213> Saccharomyces cerevisiae
<400> 120
Met Val Glu Glu Asn Ser Arg Val Leu Ile Val Leu Pro Tyr Thr Pro
 1
                  5
                                     10
Pro Ser Ala Thr Leu Gln Arg Ile Ile Gly Gln Thr Ile Pro Phe Leu
                                                     30
            20
                                 25
```

Arg	Glu	Cys 35	Gln	Ser	Gln	Leu	Asp 40	Ile	Val	Ile	Val	Pro 45	Glu	Phe	Lys
Thr	Ser 50	Phe	Gln	Leu	qaA	Ser 55	Ala	Leu	Gly	Lys	Met 60	Tyr	Ser	Ile	Thr
Arg 65	Asp	Val	Leu	Leu	Gly 70		Gly	Met	Ile	Asn 75	Ser	Gly	Ile	Asn	Ile 80
Ile	Phe	Asn	Asn	11e 85	His	Phe	Val	Glu	Ser 90	Asn	Leu	Gln	Trp	Lys 95	Val
Val	Leu	Leu	Pro 100	Gln	Glu	Ser	Thr	Phe 105	Glu	Thr	Trp	Lys	Leu 110	Glu	Leu
Gly	Gln	Gly 115	Gln	Tyr	His	Ser	11e 120	Glu	His	Tyr	Ala	Leu 125	His	Asp	Asn
Ile	Met 130	Glu	Glu	Ile	Glu	Gly 135	Pro	Lys	Asp	Ala	Asn 140	Lys	Phe	His	Val
Thr 145	Ala	Leu	Gly	Gly	Thr 150	Phe	qaA	His	Ile	His 155	Asp	Gly	His	Lys	11e
Leu	Leu	Ser	Val	Ser 165	Thr	Phe	Ile	Thr	Ser 170	Gln	Arg	Leu	Ile	Cys 175	Gly
Ile	Thr	Cys	Asp 180	Glu	Leu	Leu	Gln	Asn 185	Lys	Lys	Tyr	Lys	Glu 190	Leu	Ile
Glu	Pro	Туг 195	Asp	Thr	Arg	Cys	Arg 200	His	Val	His	Gln	Phe 205	Ile	Lys	Let
Leu	Lys 210	Pro	Asp	Leu	Ser	Val 215	Glu	Leu	Val	Pro	Leu 220	Arg	Asp	Val	Суя
Gly 225	Pro	Thr	Gly	Lys	Val 230	Pro	Glu	Ile	Glu	Суs 235	Leu	Val	Val	Ser	Arg 240
				245					250				Ile	255	
Gly	Met	Ser	Pro	Leu	Ala	Val	His	Val		Asn	Val	Leu	Gly	Gly	Arç

Glu Glu Asp Gly Trp Ser Glu Lys Leu Ser Ser Thr Glu Ile Arg Arg

285

```
Leu Leu Lys Ser Ser Ala Ser Pro Thr Cys Thr Pro Gln Asn Pro Cys
                       295
                                            300
Val
305
<210> 121
<211> 1433
<212> DNA
<213> Saccharomyces cerevisiae
<400> 121
agagagecat cegtaactet getaaggaag etgactaett tggtgatget gacaaggeca 60
ccacgattga cgaacaagtt ggtttgatcg ttgacagttt gaatgacgaa gagttagtgt 120
ccaccgccga taagatcaag gccaatgctg ctggtgccaa ggaagttttg aaggaatctg 180
caaagactat tgtcgattct ggcaaactac catccagctt gttgtcctac ttcgtgtgaa 240
taccytaaga aatggaatag aatatatacg aatgtatacg aatattatag agaacyttct 300
cttttatttc tataatgaat aggttcgggt aacggttccc tttttaggta tttctagaag 360
atgagagaag agggaataat gagaaaggcg aaaaataaag gacaccttta acgaaagatc 420
aaaggtgtcc ttatttactt acaatagctg caattagtac gactcaaaaa aagtgaaaac 480
aaaactgaaa ggatagatca atgtcttaca gaggacctat tggaaatttt ggcggtatgc 540
caatgtcatc atcgcaagga ccatactctg gcggtgcaca attcagatca aaccagaacc 600
aatccacttc tggcatctta aagcaatgga agcattcttt tgaaaagttt gcctccagaa 660
ttgaggggct cactgacaat gcagttgttt ataaattgaa gccttacatt ccaagtttgt 720
caagattttt cattgtggcc accttttatg aagattcgtt taggatctta tcacaatggt 780
cagatcaaat tttttatctg aataagtgga agcattaccc atacttcttt gtcgttgtgt 840
ttctagtggt tgttaccgtt tccatgttga ttggcgccag tttgttagtt ttaagaaagc 900
aaaccaatta tgccaccggt gtgttatgtg cttgcgttat ttctcaagca ttagtttatg 960
ggttgtttac gggttcatca tttgtcctaa gaaactttag tgttattggt gggttgttaa 1020
ttgcattcag cgattcaatt gttcaaaaca agacaacatt cggtatgctt cctgaattaa 1080
acagcaaaaa cgacaaagcg aagggttacc tgttgtttgc tggtagaatt ttaattgttt 1140
taatgtttat cgctttcact ttcagtaaat catggtttac tgttgttttg accattatcg 1200
gcacaatatg tttcgccatt ggttacaaga caaaattcgc atccattatg ttgggtttga 1260
tactaacttt ttacaatatc acgctaaaca actactggtt ttataacaat actaagagag 1320
atttcttgaa gtatgagttt taccagaact taagcatcat tggtgggctt ctattagtta 1380
ctaatactgg cgctggtgaa ttatccgttg atgaaaagaa gaagatttac tag
                                                                  1433
<210> 122
<211> 310
<212> PRT
<213> Saccharomyces cerevisiae
<400> 122
Met Ser Tyr Arg Gly Pro Ile Gly Asn Phe Gly Gly Met Pro Met Ser
```

10

15

Ser Ser Gln Gly Pro Tyr Ser Gly Gly Ala Gln Phe Arg Ser Asn Gln 20 25 30

Asn Gln Ser Thr Ser Gly Ile Leu Lys Gln Trp Lys His Ser Phe Glu 35 40 45

Lys Phe Ala Ser Arg Ile Glu Gly Leu Thr Asp Asn Ala Val Val Tyr 50 55 60

Lys Leu Lys Pro Tyr Ile Pro Ser Leu Ser Arg Phe Phe Ile Val Ala 65 70 75 80

Thr Phe Tyr Glu Asp Ser Phe Arg Ile Leu Ser Gln Trp Ser Asp Gln
85 90 95

Ile Phe Tyr Leu Asn Lys Trp Lys His Tyr Pro Tyr Phe Phe Val Val
100 105 110

Val Phe Leu Val Val Val Thr Val Ser Met Leu Ile Gly Ala Ser Leu 115 120 125

Leu Val Leu Arg Lys Gln Thr Asn Tyr Ala Thr Gly Val Leu Cys Ala 130 135 140

Cys Val Ile Ser Gln Ala Leu Val Tyr Gly Leu Phe Thr Gly Ser Ser 145 150 155 160

Phe Val Leu Arg Asn Phe Ser Val Ile Gly Gly Leu Leu Ile Ala Phe 165 170 175

Ser Asp Ser Ile Val Gln Asn Lys Thr Thr Phe Gly Met Leu Pro Glu 180 185 190

Leu Asn Ser Lys Asn Asp Lys Ala Lys Gly Tyr Leu Leu Phe Ala Gly
195 200 205

Arg Ile Leu Ile Val Leu Met Phe Ile Ala Phe Thr Phe Ser Lys Ser 210 215 220

Trp Phe Thr Val Val Leu Thr Ile Ile Gly Thr Ile Cys Phe Ala Ile 225 230 235 240

Gly Tyr Lys Thr Lys Phe Ala Ser Ile Met Leu Gly Leu Ile Leu Thr 245 250 255

Phe Tyr Asn Ile Thr Leu Asn Asn Tyr Trp Phe Tyr Asn Asn Thr Lys

260 265 270

Arg Asp Phe Leu Lys Tyr Glu Phe Tyr Gln Asn Leu Ser Ile Ile Gly 275 280 285

Gly Leu Leu Val Thr Asn Thr Gly Ala Gly Glu Leu Ser Val Asp 290 295 300

Glu Lys Lys Lys Ile Tyr 305 310

<210> 123

<211> 1802

<212> DNA

<213> Saccharomyces cerevisiae

<400> 123

catttagttc tgattacagc agaaatcgta gcgcgatgag acatttcatc aaatggcctt 60 ttttttttgg gcaatttttt tatatcttga aatgatagtt gccttgtact ttcaaccgtt 120 catttcatta agaacttgac taaatatgaa catttcttaa aaaaaaggtt gacatataaa 180 aactataaat atgtcaaaga aaccatacaa tcatagattt ataactatct tttggatgac 300 attaatgaac ataacgctcc taatacaaat gtccaaaaaa tattacccgc aaatacgaat 360 ctttttttt tctcgatgaa attttgcaaa gagttcgaaa tttttatttc aagagctggt 420 agagaaaatt tcataaggtt ttcctaccga tgcttttata aaatcttcgt tttgtctcac 480 atataccaac aagagtaacg atgttttctt tacctaccct aacctcagac atcactgttg 540 aagtcaacag ttccgctacc aaaaccccat tcgtccgtcg tccggtcgaa ccggttggta 600 agttcttttt gcaacatgct caaagaactt tgagaaacca cacctggtct gaatttgaaa 660 gaattgaagc tgaaaagaac gtcaaaaccg ttgatgaatc caatgtcgac ccagatgagt 720 tgttattcga cactgaattg gccgatgaag atttactgac tcatgatgct agagactgga 780 aaactgccga tttgtatgct gctatgggtt tgtctaagtt gcgtttcaga gctactgaaa 840 gtcaaatcat caaggctcac agaaaacaag ttgtcaagta ccatccagac aagcaatctg 900 ctgctggtgg tagtttggac caagatggct ttttcaagat tattcaaaag gcctttgaaa 960 ctttgactga ttccaacaag agagctcagt acgactcatg tgattttgtt gccgatgttc 1020 ctcctccaaa gaagggtacc gattatgact tttatgaagc ttggggcccc gttttcgaag 1080 ctgaagctcg tttttctaag aagactccta ttccttctct aggtaacaaa gattcttcca 1140 agaaggaagt tgaacaattc tatgctttct ggcacagatt tgactcctgg agaacctttg 1200 agttcttgga cgaagatgtc ccagatgact cttctaacag agaccacaag cgttacattg 1260 aaagaaagaa caaggccgca agagacaaga agaagactgc tgataacgct agattggtca 1320 aacttgttga aagagctgtc agtgaagatc cccgtatcaa aatgttcaaa gaagaagaga 1380 agaaggaaaa ggaaagaaga aaatgggaaa gagaagccgg tgccagagct gaagctgaag 1440 ctaaggccaa ggccgaagct gaagcgaagg ctaaagctga atctgaagcc aaggctaacg 1500 cctccgcaaa agctgacaaa aagaaggcta aggaagctgc taaggccgcc aagaaaaaga 1560 acaagagagc catccgtaac tctgctaagg aagctgacta ctttggtgat gctgacaagg 1620 ccaccacgat tgacgaacaa gttggtttga tcgttgacag tttgaatgac gaagagttag 1680 tgtccaccgc cgataagatc aaggccaatg ctgctggtgc caaggaagtt ttgaaggaat 1740

ctgcaaagac tattgtcgat tctggcaaac taccatccag cttgttgtcc tacttcgtgt 1800 ga

<210> 124

<211> 433

<212> PRT

<213> Saccharomyces cerevisiae

<400> 124

Met Phe Ser Leu Pro Thr Leu Thr Ser Asp Ile Thr Val Glu Val Asn 1 5 10 15

Ser Ser Ala Thr Lys Thr Pro Phe Val Arg Arg Pro Val Glu Pro Val 20 25 30

Gly Lys Phe Phe Leu Gln His Ala Gln Arg Thr Leu Arg Asn His Thr 35 40 45

Trp Ser Glu Phe Glu Arg Ile Glu Ala Glu Lys Asn Val Lys Thr Val 50 55 60

Asp Glu Ser Asn Val Asp Pro Asp Glu Leu Leu Phe Asp Thr Glu Leu 65 70 75 80

Ala Asp Glu Asp Leu Leu Thr His Asp Ala Arg Asp Trp Lys Thr Ala 85 90 95

Asp Leu Tyr Ala Ala Met Gly Leu Ser Lys Leu Arg Phe Arg Ala Thr
100 105 110

Glu Ser Gln Ile Ile Lys Ala His Arg Lys Gln Val Val Lys Tyr His 115 120 125

Pro Asp Lys Gln Ser Ala Ala Gly Gly Ser Leu Asp Gln Asp Gly Phe 130 135 140

Phe Lys Ile Ile Gln Lys Ala Phe Glu Thr Leu Thr Asp Ser Asn Lys 145 150 155 160

Arg Ala Gln Tyr Asp Ser Cys Asp Phe Val Ala Asp Val Pro Pro Pro 165 170 175

Lys Lys Gly Thr Asp Tyr Asp Phe Tyr Glu Ala Trp Gly Pro Val Phe 180 185 190

Glu Ala Glu Ala Arg Phe Ser Lys Lys Thr Pro Ile Pro Ser Leu Gly
195 200 205

Asn	Lys	Asp	Ser	Ser	Lys	Lys	Glu	Val	GLu	GIn	Pne	īĀī	Ala	Pne	тр
	210					215					220				

His Arg Phe Asp Ser Trp Arg Thr Phe Glu Phe Leu Asp Glu Asp Val 225 230 235 240

Pro Asp Asp Ser Ser Asn Arg Asp His Lys Arg Tyr Ile Glu Arg Lys 245 250 255

Asn Lys Ala Ala Arg Asp Lys Lys Lys Thr Ala Asp Asn Ala Arg Leu 260 265 270

Val Lys Leu Val Glu Arg Ala Val Ser Glu Asp Pro Arg Ile Lys Met 275 280 285

Phe Lys Glu Glu Glu Lys Lys Glu Lys Glu Arg Arg Lys Trp Glu Arg 290 295 300

Glu Ala Gly Ala Arg Ala Glu Ala Glu Ala Lys Ala Lys Ala Glu Ala 305 310 315 320

Glu Ala Lys Ala Lys Ala Glu Ser Glu Ala Lys Ala Asn Ala Ser Ala 325 330 335

Lys Ala Asp Lys Lys Lys Ala Lys Glu Ala Ala Lys Ala Ala Lys Lys 340 345 350

Lys Asn Lys Arg Ala Ile Arg Asn Ser Ala Lys Glu Ala Asp Tyr Phe 355 360 365

Gly Asp Ala Asp Lys Ala Thr Thr Ile Asp Glu Gln Val Gly Leu Ile 370 375 380

Val Asp Ser Leu Asn Asp Glu Glu Leu Val Ser Thr Ala Asp Lys Ile 385 390 395 400

Lys Ala Asn Ala Ala Gly Ala Lys Glu Val Leu Lys Glu Ser Ala Lys
405 410 415

Thr Ile Val Asp Ser Gly Lys Leu Pro Ser Ser Leu Leu Ser Tyr Phe 420 425 430

Val

```
<210> 125
<211> 1472
 <212> DNA
<213> Saccharomyces cerevisiae
<400> 125
gacteggaeg gaacegtete atattataeg tteagtgaat attttteaeg gaagaatgga 60
aagggcctcg aaagacgtta atgcatccgt acacctacat ctttacattt tttgctgttg 120
cacctacact gtttttttt ttttcacctt atgagtcctg tatttcttga aagagccgat 180
aacaatattc caggtggagt cccgaggcag aatcaaggct gcggagagaa gttcctctca 240
aactaatggg agtgatccgc tcagttctcc catcacgaca gaactgtctg agacaaagtc 300
tttccagcag agtccgccta cgctcttgct gcagagattc gcccgaaggc aggtttcctg 360
aaattetttt cactagtaaa gtgttegtte atgtaaaaca tactgeegta gttttgaget 420
aaaattaaag atatattaga tttttagaat ttcttagata gtctcaacgt gttaaaacaa 480
aagcataacc aaagaaaaaa atggctaagt tettgaaagc tggtaaagtt ggtacgtatc 540
attttcagtt tttggacatc aacaaaaaac cgtgcaactg tggtattaga aatccaagtt 600
attagcagtt gatgttgtat aatttgagtt tgaattcgag tattgtatct tcatatggga 660
gtagggagga aaaagcgtgg agatgttaga tctgggaaaa aacctttaaa tttgaatgat 720
agcttattaa tagaaatatt ggacaaaaag atgaatgatt taggagcgaa actaacgatt 780
attcatattc attgagctat acgaggcagt ggaagggtta tcaaaaagtt gtaatttaaa 840
agaaaaaagc atcaatagcc accaaataag atggtgaaag ttccgcgata tatttgctaa 900
gaccataggc acatgtacgc aggcgaccag ggccttttat ttactgtagt tctttccatg 960
ccctattaaa aattcgtggg cctatgataa tttttctctc cctagtattg aagaacaaaa 1020
ttgagaattc gccaaacttt aagaagggtt cacatttact aactatttct tttccttcct 1080
ttttttacac agctgtcgtt gtccgtggtc gttacgccgg taagaaggtt gttatcgtta 1140
aaccacatga tgaaggttcc aagtctcacc catttggtca cgctttggtt gccggtattg 1200
aaagataccc attgaaggtc accaagaagc acggtgccaa gaaggttgct aagagaacca 1260
agatcaagcc tttcatcaag gtcgtcaact acaaccattt attgccaacc agatacactt 1320
tagatgttga agctttcaaa tccgttgttt ctactgaaac tttcgaacaa ccttcccaac 1380
gtgaagaagc taagaaagtc gtcaagaagg ctttcgaaga aagacaccaa gctggtaaga 1440
accaatggtt cttctctaag ttgagatttt aa
                                                                  1472
<210> 126
<211> 136
<212> PRT
<213> Saccharomyces cerevisiae
<400> 126
Met Ala Lys Phe Leu Lys Ala Gly Lys Val Ala Val Val Arg Gly
                  5
                                     10
Arg Tyr Ala Gly Lys Lys Val Val Ile Val Lys Pro His Asp Glu Gly
            20
                                 25
                                                     30
```

Ser Lys Ser His Pro Phe Gly His Ala Leu Val Ala Gly Ile Glu Arg

```
Tyr Pro Leu Lys Val Thr Lys Lys His Gly Ala Lys Lys Val Ala Lys
                         55
Arg Thr Lys Ile Lys Pro Phe Ile Lys Val Val Asn Tyr Asn His Leu
                                         75
                     70
Leu Pro Thr Arg Tyr Thr Leu Asp Val Glu Ala Phe Lys Ser Val Val
                                     90
                 85
Ser Thr Glu Thr Phe Glu Gln Pro Ser Gln Arg Glu Glu Ala Lys Lys
                                                    110
            100
                                105
Val Val Lys Lys Ala Phe Glu Glu Arg His Gln Ala Gly Lys Asn Gln
                                                125
        115
                            120
Trp Phe Phe Ser Lys Leu Arg Phe
                        135
    130
<210> 127
<211> 1299
<212> DNA
<213> Saccharomyces cerevisiae
<400> 127
ttacttacgt ctattataac cttccgcgta aaaaagaaat attgacatcc tttcttgtat 60
ttagcccata cattttcacc catgcaccat tggattataa agaaaaaaat ttaataaaaa 120
tctgccgggg aaatttcaga agaaaaaagg aaggtgtgtt cgcatttaac acgggccacc 180
ataaactttt gtttgccacc catctagacg ggatccgccc cgcccaaggc tctcttcctc 240
tagctaggca atgtggctct cggaaaggaa actcccacca ggacgtggtg ggaaatgcag 300
caattcccct ctgcttcccg ctgacctttc ttgggcccag tttagtaagt atgctctcat 360
ctttcatgtg tgctgtaact tgcaagtcat taacactatc tataattgat attagacgtt 420
aaaagttcta tatagcattt gcttttattt agagaatacg aaaacacacc agataattag 480
tgcatatata ttagatcaat atggtatgtg aaagagatat taaacataag atgtgaagga 540
aacagaagag aacaatattt gaggaatgaa ttcggatgaa attctagtag aagaaaaaag 600
cctaaaataa aaagaaagaa aggaaaataa agcagtaatc aacctctgac atttgatagg 660
agtatgcgtt tccgcaaaat caacaaatca tgcattaaac tggtaacgga gactttaaag 720
acgtgcaaag aaagcaatta atttttagat agggaattga aaggctctgt atagcagaaa 780
atttgagatc tactgtggag ataagcatat ggtgttgagt tatgggtaat ctaacaataa 840
tagaaactta tagcagttga agatggcgga tcatgatgcc ggatactgct gggtaatgat 900
ttatccttct ttttttggag cacaaatgct cactttttcc tgtctcctct aacttttctg 960
atgtatecta eccaacecta cagtttteaa attactaaca teaatttttt tgtgaaacga 1020
```

gccaaattat ctgaaggtac ttctttcaga agaaagtaa

ctttcgtttt cgtttctatt tatttcatta caggttttag ttcaagattt gttgcacca 1080 actgctgctt ctgaagccag aaagcacaag ttaaagactt tggttcaagg cccaagatcc 1140 tacttcttgg atgtcaaatg cccaggttgt ttgaacatca ccactgtttt ttctcatgct 1200 caaactgctg tcacttgtga atcatgctct actgtcttat gtactccaac cggtggtaag 1260

```
<210> 128
<211> 82
<212> PRT
<213> Saccharomyces cerevisiae
<400> 128
Met Val Leu Val Gln Asp Leu Leu His Pro Thr Ala Ala Ser Glu Ala
Arg Lys His Lys Leu Lys Thr Leu Val Gln Gly Pro Arg Ser Tyr Phe
             20
                                  25
Leu Asp Val Lys Cys Pro Gly Cys Leu Asn Ile Thr Thr Val Phe Ser
         35
                              40
                                                  45
His Ala Gln Thr Ala Val Thr Cys Glu Ser Cys Ser Thr Val Leu Cys
     50
                         55
Thr Pro Thr Gly Gly Lys Ala Lys Leu Ser Glu Gly Thr Ser Phe Arg
                     70
                                          75
Arg Lys
<210> 129
<211> 1262
<212> DNA
<213> Saccharomyces cerevisiae
<400> 129
taaaacatgg cattttaaat agtactgcca aaatttaatg caatggcaat tagttggcta 60
tgttgctaca tattattttc catagcattc tctgtcacaa acttcaaaat aaaagacagt 120
gacateegaa cateegatae acceataeat catgaaaata aateetgaca ttggaagaag 180
tcttgataca aaaaaagatc gctgtcaaaa atgcggccag actgccgctg catcgtacca 240
acagtgcgta tgttcagacg gagagacgac ctctagagag acgtccgtcg gtgcggcaac 300
gctgacggtt tagttgttcg acgggatgat gggttccgcc agggggaggg aaggctttcc 360
accaagagag gtaaaattat tcgtcgaaat gaactcagag atacatccat attgttgaca 420
atgtatatet taattgatgt ggtattttea etgttttaae gtaaattgaa ggagattaag 480
caaaaaaaca atcagtaata atgggtatgt ggacgattag gaatagacaa accatgttat 540
ttatctccat tagggcgtga gagtgtaatt agtacacagg tactactaga atgctaaaga 600
```

actttttaaa atatcctgaa tcgtagggca aatccatgtc aagcaagaaa ctaatagtta 660 ttaaacttca tttacttttg agctagttaa atattttcat catttcctaa agtactgaac 720 acctgaatga tacttttatt ggccctttta ataagaactc tggttagaaa atatattgag 780 gatatcatta gtaatactca ttagatattt gtgaatttag ccgtttcccc attacagaaa 840

```
aaagatacaa ctaattacat gtgcagtcaa attacttttt ttttaagatc aattactaac 900
aatcaactat catgctaaat ttgctgtgat atcattttga accagttaac gtcccaaaga 960
ccagaaagac ctactgtaag ggtaagacct gtcgtaagca cactcaacac aaggttactc 1020
aatacaaagc tggtaaggct tccttgttcg ctcaaggtaa gagacgttat gaccgtaaac 1080
aatctggttt cggtggtcaa accaagcctg ttttccacaa gaaagctaag actaccaaga 1140
aggttgtttt gagattggaa tgtgtcaaat gtaagactag agcccaatta accttgaaga 1200
gatgtaagca cttcgaattg ggtggtgaaa agaagcaaaa gggtcaagct ttgcaattct 1260
                                                                   1262
ga
<210> 130
<211> 106
<212> PRT
<213> Saccharomyces cerevisiae
<400> 130
Met Val Asn Val Pro Lys Thr Arg Lys Thr Tyr Cys Lys Gly Lys Thr
                  5
                                     10
Cys Arg Lys His Thr Gln His Lys Val Thr Gln Tyr Lys Ala Gly Lys
                                 25
Ala Ser Leu Phe Ala Gln Gly Lys Arg Arg Tyr Asp Arg Lys Gln Ser
                             40
Gly Phe Gly Gly Gln Thr Lys Pro Val Phe His Lys Lys Ala Lys Thr
     50
                         55
                                             60
Thr Lys Lys Val Val Leu Arg Leu Glu Cys Val Lys Cys Lys Thr Arg
 65
                     70
                                         75
                                                              80
Ala Gln Leu Thr Leu Lys Arg Cys Lys His Phe Glu Leu Gly Gly Glu
                 85
Lys Lys Gln Lys Gly Gln Ala Leu Gln Phe
            100
                                105
<210> 131
<211> 962
<212> DNA
<213> Saccharomyces cerevisiae
<400> 131
catcgcttga tttccggcct gcaaaaataa agtagtcggt acgtactttc gttttcaatt 60
```

tccatggtgc acagtatctt aactatctgc ttagtcgagg agaaccagga ttctgttcgt 120 tgctcagccg cttcgtggat attctcttgg atactttaaa catggaccta cgttccgctc 180

02-07-1999

<210> 132

<211> .153

<212> PRT

<213> Saccharomyces cerevisiae

<400> 132

Met Ser Leu Arg Pro Cys Leu Thr Pro Ser Ser Met Gln Tyr Ser Asp 1 5 10 15

Ile Tyr Ile His Thr Pro His Pro His Pro His Pro His Pro His Thr
20 25 30

Pro Thr His Thr His Pro His Thr Pro Thr Pro Thr Pro His Pro His 35

Pro His Thr Pro His Pro His Thr Thr Pro Thr Pro Thr Pro His His 50 55 60

Thr His Thr Pro His Thr Thr Leu Ser Asn Leu Ser Leu Asn Leu Pro 65 70 75 80

Ser His Tyr Pro Thr Ser Pro Leu Val Thr Leu Pro His Ser Thr Ile 85 90 95

Pro Leu Pro Thr Thr Ile His Leu Ser Thr Tyr Tyr Tyr His Pro Pro 100 105 110

Pro Ile Ile Thr Val Thr Leu Gln Leu Pro Ile Ser Asn Ser Thr Thr 115 120 125

Ile Thr Leu Leu Pro Tyr His Pro Pro Cys Pro Thr His Cys Thr 130 135 140 Val Val Leu Pro Ser Ile Leu Lys Arg 145 150

<210> 133 <211> 3752 <212> DNA <213> Saccharomyces cerevisiae

#### <400> 133

ttccataggg cttattttcc agttgatgga atgggaggtt gctcttaacg cgaagactaa 60 ttcaatgagg ctttattgag gggcacgatg ctctttttt cttctgtact ttataaataa 180 ttccgtattt cttcgctttg tttcatgctc tgttctgagc ttttagtttc tcttattgag 240 agccttttcc aaatcaaggc tcaggaagtg actcgtcctt agaagaacaa ggtttggatt 300 tggtagggtt cttgtacgaa gttctcaata taatttgcgc atttactgta cggttcatac 360 tagtttattc cgggtaacaa gttttcttgt gatgctaaat caatgtgtat attgagaaaa 420 ctatgtagta agttacacaa agcaacaaag gatattatta tatgtgacag agaagaattg 480 ctgtagagat tcatgacaat atgaatgaga ctactacaaa acagccgcta aagaaaaggt 540 ccctgagcag ctatctttca aacgtgagca cgagacggga ggagctggaa aagatttcta 600 aacaagaaac atccgaagag gaagatactg ccggcaagca tgaacaaagg gaaacactgt 660 cggaagaagt aagtgataag tttccagaaa atgttgcatc gtttcgctca cagactacaa 720 gegtteatea agecacecaa aataacetga atgeaaagga ateegaagae etggeecata 780 agaatgatgc gagttcacac gaaggagagg ttaatgggga cagccgtccg gacgatgttc 840 ctgaaacgaa tgaaaaaata agccaagcaa tacgagcgaa aatttcgtca tcatcatcat 900 cacccaatgt acgtaatgtt gatatccaaa accatcaacc tttttcgaga gaccaacttc 960 gagcgatgtt gaaagaacca aaaaggaaaa ctgttgatga tttcatagaa gaagagggtt 1020 tgggagctgt tgaagaagag gatttaagtg atgaggtact tgaaaaaaat acaacagaac 1080 cagaaaatgt ggaaaaagat atagagtata gcgactcaga taaggacaca gacgatgtgg 1140 gaagcgatga tcccacggca cccaactcac caataaaact tggtcgtcgc aaactggtta 1200 gaggtgacca acttgatgca acaacaagtt ccatgtttaa taacgaatca gattctgaat 1260 tatcagatat cgatgatagc aagaatattg ctttatccag tagcctattt agaggcggtt 1320 cttcacctgt gaaagagaca aacaacaacc tttcaaatat gaattcttca ccagcacaaa 1380 atccaaaaag gggttctgtc tccaggagta atgatagtaa caagagttct catatagctg 1440 tttccaagcg ccccaaacag aagaagggca tatataggga ttctggtggt agaacaagac 1500 tacagattgc ctgtgacaag ggcaaatatg atgtagtcaa aaagatgatt gaagaaggag 1560 gttacgatat taatgaccaa gacaatgctg gtaatacagc attgcatgaa gcggcgttgc 1620 aaggtcatat tgagattgtg gaactgttga tagaaaatgg tgcagatgta aatatcaagt 1680 ctattgaaat gttcggcgat actcccttga tcgatgcttc cgccaatgga cacttggatg 1740 ttgtcaagta tcttcttaaa aacggtgcgg acccaactat acgtaacgct aaagggttaa 1800 ctgcgtttga atctgtcgat gatgaatctg aatttgatga tgaagaagac caaaagattt 1860 tgcgtgaaat aaaaaaaagg ttgagtatag ccgctaaaaa atggactaac agagcaggaa 1920 ttcataatga caaatctaaa aatggcaata atgctcacac aatagatcag ccaccttttg 1980 ataataccac aaaagccaaa aacgaaaagg ccgctgactc accttcaatg gcttccaata 2040 ttgatgagaa agctccggaa gaggaattct attggacaga tgttacttct agagcgggaa 2100 aagaaaaact gtttaaagct tcgaaggagg gacatttacc atacgttggt acgtatgtag 2160

Printed:17-08-2000 151

```
aaaatggtgg taagatagac ttaaggtcat ttttcgaaag cgttaagtgt ggccatgaag 2220
atattacaag tatctttttg gcattcggat ttcccgtaaa tcaaacttca agggataata 2280
aaacatccgc tttaatggta gctgtaggtc gtggccatct tggaactgtt aaactgctat 2340
tagaggcagg tgcagatcca accaaaagag ataaaaaggg tcgtaccgct ttgtactatg 2400
ccaaaaacag cataatggga ataacaaaca gtgaagaaat tcaattgatt gaaaatgcta 2460
taaataacta tttgaagaag cactctgaag ataataatga tgatgatgat gatgatgata 2520
ataataatga aacatataaa catgaaaaaa agagagagaa aacgcaatca cccatattag 2580
caagccgaag aagtgccact cctagaattg aagacgaaga ggacgatacg aggatgctca 2640
atctggcaga cgatgacttt aataacgatc gtgatgtcaa ggaatctaca acttcggatt 2700
cgcgcaagag attggatgac aatgaaaatg ttggtactca atactcattg gactggaaaa 2760
aacgtaaaac aaatgccttg caagatgaag aaaaattaaa aagtatctca ccactctcta 2820
tggaacctca ttctccaaag aaggcaaaat ctgtagagat aagtaaaata catgaagaaa 2880
cggctgctga aagagaagca agactcaagg aagaggaaga atacagaaag aaaagattag 2940
aaaagaaaag aaaaaaagaa caggaactac tacaaaagct ggccgaagat gagaaaaaaa 3000
ggatcgaaga acaggagaag cagaaagtct tagaaatgga aagattggaa aaagctactt 3060
tagagaaagc aagaaaatg gaaagggaaa aagaaatgga agaaatctct tatagaaggg 3120
cagtaaggga cttatatccg ttaggactga agattattaa cttcaacgat aaacttgatt 3180
acaaaagatt tttgccgcta tattattttg tagacgaaaa aaacgataaa tttgtgctcg 3240
acttgcaagt aatgatattg ttgaaggata tcgacttgct ctcaaaggac aaccaaccaa 3300
cttctgagaa gattcccgtc gatccctcgc acctgactcc attgtggaat atgttgaaat 3360
tcattttcct gtatggaggt agttatgatg ataaaaagaa caacatggaa aataaaagat 3420
atgttgtaaa ctttgatggg gttgatttgg acacaaagat tgggtatgag cttttggagt 3480
acaaaaaatt tgttagtttg cccatggcgt ggattaaatg ggataacgtt gttattgaaa 3540
accacgcaaa aagaaaagaa attgaaggaa atatgattca aatatcaata aacgaatttg 3600
cacgatggag aaacgataag ctgaacaaag cgcagcagcc gacgcgcaaa cagcgttctc 3660
taaaaatacc tcgagagtta ccggttaaat ttcaacaccg tatgagcata tcctccgtcc 3720
tccagcagac atccaaagaa ccattttggt aa
                                                                  3752
```

```
<210> 134
```

<211> 1083

<212> PRT

<213> Saccharomyces cerevisiae

### <400> 134

Met Asn Glu Thr Thr Lys Gln Pro Leu Lys Lys Arg Ser Leu Ser

1 5 10 15

Ser Tyr Leu Ser Asn Val Ser Thr Arg Arg Glu Glu Leu Glu Lys Ile 20 25 30

Ser Lys Gln Glu Thr Ser Glu Glu Glu Asp Thr Ala Gly Lys His Glu
35 40 45

Gln Arg Glu Thr Leu Ser Glu Glu Val Ser Asp Lys Phe Pro Glu Asn 50 55 60

Val Ala Ser Phe Arg Ser Gln Thr Thr Ser Val His Gln Ala Thr Gln

65					70					75					80
Asn	Asn	Leu	Asn	Ala 85	Lys	Glu	Ser	Glu	Asp 90	Leu	Ala	His	Lys	Asn 95	Asp
Ala	Ser	Ser	His 100	Glu	Gly	Glu	Val	Asn 105	Gly	Asp	Ser	Arg	Pro 110	Asp	Asp
Va1	Pro	Glu 115	Thr	Asn	Glu	Lys	Ile 120	Ser	Gln	Ala	Ile	Arg 125	Ala	Lys	Ile
Ser	Ser 130	Ser	Ser	Ser	Ser	Pro 135	Asn	Val	Arg	Asn	Val 140	Asp	Ile	Gln	Asn
His 145		Pro	Phe	Ser	Arg 150	Asp	Gln	Leu	Arg	Ala 155	Met	Leu	Lys	Glu	Pro 160
Lys	Arg	Lys	Thr	Val 165		Asp	Phe	Ile	Glu 170	Glu	Glu	Gly	Leu	Gly 175	Ala
Val	. Glu	Glu	Glu 180		Leu	Ser	Asp	Glu 185	Val	Leu	Glu	Lys	190	Thr	Thr
Glu	ı Pro	Glu 195		Val	Glu	Lys	Asp 200		: Glu	туг	Ser	205	Ser	Asp	) Lys
As <u>r</u>	210		) Asp	Val	Gly	Ser 215		Asp	Pro	Thi	220	a Pro	) Asr	ı Sei	r Pro
11e 225		s Lev	ı Gly	Arg	230	Lys	Leu	val	L Arg	g Gly 23!	y Ası	Glı	n Lev	ı Ası	240
<b>ጥ</b> ኪ ነ	· r Thi	c Sei	s Ser	Met	: Phe	a Asn	a Asr	ı Glı	ı Sei	r Asj	o Se	r Gl	u Le	u Se	r Ası

Thr Thr Ser Ser Met Phe Asn Asn Glu Ser Asp Ser Glu Leu Ser Asp 245 250 255

Ile Asp Asp Ser Lys Asn Ile Ala Leu Ser Ser Ser Leu Phe Arg Gly 260 265 270

Gly Ser Ser Pro Val Lys Glu Thr Asn Asn Leu Ser Asn Met Asn 275 280 285

Ser Ser Pro Ala Gln Asn Pro Lys Arg Gly Ser Val Ser Arg Ser Asn 290 295 300

Asp Ser Asn Lys Ser Ser His Ile Ala Val Ser Lys Arg Pro Lys Gln 305 310 315 320

Lys Lys Gly Ile Tyr Arg Asp Ser Gly Gly Arg Thr Arg Leu Gln Ile

330

- Ala Cys Asp Lys Gly Lys Tyr Asp Val Val Lys Lys Met Ile Glu Glu 340 345 350
- Gly Gly Tyr Asp Ile Asn Asp Gln Asp Asn Ala Gly Asn Thr Ala Leu 355 360 365
- His Glu Ala Ala Leu Gln Gly His Ile Glu Ile Val Glu Leu Leu Ile 370 375 380
- Glu Asn Gly Ala Asp Val Asn Ile Lys Ser Ile Glu Met Phe Gly Asp 385 390 395 400
- Thr Pro Leu Ile Asp Ala Ser Ala Asn Gly His Leu Asp Val Val Lys
  405 410 415
- Tyr Leu Leu Lys Asn Gly Ala Asp Pro Thr Ile Arg Asn Ala Lys Gly
  420 425 430
- Leu Thr Ala Phe Glu Ser Val Asp Asp Glu Ser Glu Phe Asp Asp Glu
  435 440 445
- Glu Asp Gln Lys Ile Leu Arg Glu Ile Lys Lys Arg Leu Ser Ile Ala 450 455 460
- Ala Lys Lys Trp Thr Asn Arg Ala Gly Ile His Asn Asp Lys Ser Lys 465 470 475 480
- Asn Gly Asn Asn Ala His Thr Ile Asp Gln Pro Pro Phe Asp Asn Thr
  485 490 495
- Thr Lys Ala Lys Asn Glu Lys Ala Ala Asp Ser Pro Ser Met Ala Ser 500 505 510
- Asn Ile Asp Glu Lys Ala Pro Glu Glu Glu Phe Tyr Trp Thr Asp Val 515 520 525
- Thr Ser Arg Ala Gly Lys Glu Lys Leu Phe Lys Ala Ser Lys Glu Gly 530 535 540
- His Leu Pro Tyr Val Gly Thr Tyr Val Glu Asn Gly Gly Lys Ile Asp 545 550 555 560
- Leu Arg Ser Phe Phe Glu Ser Val Lys Cys Gly His Glu Asp Ile Thr 565 570 575
- Ser Ile Phe Leu Ala Phe Gly Phe Pro Val Asn Gln Thr Ser Arg Asp

585

590

Asn Lys Thr Ser Ala Leu Met Val Ala Val Gly Arg Gly His Leu Gly 595 600 605

Thr Val Lys Leu Leu Glu Ala Gly Ala Asp Pro Thr Lys Arg Asp 610 615 620

Lys Lys Gly Arg Thr Ala Leu Tyr Tyr Ala Lys Asn Ser Ile Met Gly 625 630 635 640

Ile Thr Asn Ser Glu Glu Ile Gln Leu Ile Glu Asn Ala Ile Asn Asn 645 650 655

Tyr Leu Lys Lys His Ser Glu Asp Asn Asn Asp Asp Asp Asp Asp Asp 660 665 670

Asp Asn Asn Glu Thr Tyr Lys His Glu Lys Lys Arg Glu Lys Thr 675 680 685

Gln Ser Pro Ile Leu Ala Ser Arg Arg Ser Ala Thr Pro Arg Ile Glu 690 695 700

Asp Glu Glu Asp Asp Thr Arg Met Leu Asn Leu Ala Asp Asp Asp Phe 705 710 715 720

Asn Asn Asp Arg Asp Val Lys Glu Ser Thr Thr Ser Asp Ser Arg Lys
725 730 735

Arg Leu Asp Asp Asn Glu Asn Val Gly Thr Gln Tyr Ser Leu Asp Trp 740 745 750

Lys Lys Arg Lys Thr Asn Ala Leu Gln Asp Glu Glu Lys Leu Lys Ser 755 760 765

Ile Ser Pro Leu Ser Met Glu Pro His Ser Pro Lys Lys Ala Lys Ser 770 780

Val Glu Ile Ser Lys Ile His Glu Glu Thr Ala Ala Glu Arg Glu Ala 785 790 795 800

Arg Leu Lys Glu Glu Glu Glu Tyr Arg Lys Lys Arg Leu Glu Lys Lys 805 810 815

Arg Lys Lys Glu Gln Glu Leu Leu Gln Lys Leu Ala Glu Asp Glu Lys 820 825 830

Lys Arg Ile Glu Glu Glu Lys Gln Lys Val Leu Glu Met Glu Arg

840

845

Leu Glu Lys Ala Thr Leu Glu Lys Ala Arg Lys Met Glu Arg Glu Lys 850 855 860

Glu Met Glu Glu Ile Ser Tyr Arg Arg Ala Val Arg Asp Leu Tyr Pro 865 870 875 880

Leu Gly Leu Lys Ile Ile Asn Phe Asn Asp Lys Leu Asp Tyr Lys Arg 885 890 895

Phe Leu Pro Leu Tyr Tyr Phe Val Asp Glu Lys Asn Asp Lys Phe Val 900 905 910

Leu Asp Leu Gln Val Met Ile Leu Leu Lys Asp Ile Asp Leu Leu Ser 915 920 925

Lys Asp Asn Gln Pro Thr Ser Glu Lys Ile Pro Val Asp Pro Ser His 930 935 940

Leu Thr Pro Leu Trp Asn Met Leu Lys Phe Ile Phe Leu Tyr Gly Gly 945 950 955 960

Ser Tyr Asp Asp Lys Lys Asn Asn Met Glu Asn Lys Arg Tyr Val Val 965 970 975

Asn Phe Asp Gly Val Asp Leu Asp Thr Lys Ile Gly Tyr Glu Leu Leu 980 985 990

Glu Tyr Lys Lys Phe Val Ser Leu Pro Met Ala Trp Ile Lys Trp Asp 995 1000 1005

Asn Val Val Ile Glu Asn His Ala Lys Arg Lys Glu Ile Glu Gly Asn 1010 1015 1020

Met Ile Gln Ile Ser Ile Asn Glu Phe Ala Arg Trp Arg Asn Asp Lys 1025 1030 1035 1040

Leu Asn Lys Ala Gln Gln Pro Thr Arg Lys Gln Arg Ser Leu Lys Ile 1045 1050 1055

Pro Arg Glu Leu Pro Val Lys Phe Gln His Arg Met Ser Ile Ser Ser 1060 1065 1070

Val Leu Gln Gln Thr Ser Lys Glu Pro Phe Trp 1075 1080

```
<210> 135
<211> 4883
<212> DNA
<213> Saccharomyces cerevisiae
```

# <400> 135 tttcaatgcg ctacaactga ctgaaccagt ccacaaattt ggctggtctc tatcgttctc 60 gccctgagag attttatttt cttataaatt ttactaaata aggatttgta cttttgatag 120 agtttttttt ttttctgacg tttgtttagt taatattata taatagtatt taggaaacta 180 ggtgggaggg gacatcgcaa cttttatcgt gacctacgac cagtgttttt tcaattgttt 240 gatcaaaaga gaaaaaaaga caaaggaccg aaggatagtt gcacgcataa actggagaaa 300 aaattgtgtt tgacatcggc gtttaggctt gtttgttctg tcacacatac gctgcttcac 360 accaattcat atttctcagg ttaatttgtc tcctctccaa cttcaataac gattttgcgt 420 gaaggtttaa tttcattatt gcaattttag cagagaaacg cacataaata tatatatt 480 gattacagaa ccattataac atgtcttctt tgaaggatga agtacccact gagacttccg 540 aagacttcgg ttttaagttt ttaggtcaaa aacaaattct accttccttc aatgaaaaac 600 tgccatttgc atctctacaa aatctcgata tttcaaacag taagtcttta ttcgttgctg 660 cctctggtag taaggcggtg gtcggcgaat tacaattact gagagatcat atcacctccg 720 actictactic gitaacgite aagtgggaga aagaaatice agatgtaata titgtgtgct 780 ttcatggtga tcaggttttg gtttcaacca gaaatgcatt atattcgtta gacttggagg 840 aattgagtga atttcgaacg gtcacttctt ttgagaagcc agttttccaa ttgaagaacg 900 ttaataacac tttagtaatt ttaaattcag tcaatgattt atcagcactg gatttaagaa 960 caaaatcgac taagcaactg gcacaaaacg ttacctcttt tgatgtcaca aattcgcagt 1020 tagcagttct actaaaagat agaagttttc aaagttttgc atggcgaaat ggcgaaatgg 1080 aaaaacaatt tgagttctct ctaccgtcag aattagaaga gcttccagta gaagaatatt 1140 cccctttgag tgttaccatt ctctctccac aggatttttt ggcggttttc ggtaatgtta 1200 tatcagagac cgatgacgaa gtttcatacg atcaaaaaat gtacattata aagcacatag 1260 acggcagcgc ctcatttcaa gaaacttttg atattacacc tccattcggg caaatagtaa 1320 ggttcccata tatgtacaaa gttaccttgt ctggtttaat tgaacctgat gcaaacgtaa 1380 atgtgctagc atcatcatgt tcaagtgaag taagtatatg ggactcgaaa caagttattg 1440 aaccttccca ggattctgaa cgagcagtat tgcccatcag tgaggaaaca gataaggaca 1500 caaatccaat aggtgtggca gttgacgtcg ttacttcagg cactattcta gaaccttgtt 1560 ccggtgttga tacgatagag cgattgccgc tcgtttacat attgaataac gaaggtagct 1620 tacagatagt cgggttgttt catgtggcag caatcaaaag cggccattat agcataaatc 1680 tggaatettt agaacatgag aaatetetet etectacate agaaaaaatt cetattgetg 1740 gacaggagca ggaagaaaaa aagaaaaata atgaatcaag taaggcttta tcagagaatc 1800 ctttcacatc agcaaataca tcaggcttca cttttcttaa aacacaacca gccgctgcca 1860 atagcctgca gtctcaaagt tcttcaacct ttggtgctcc ctcatttgga tcatccgcat 1920 ttaaaattga cttgccatca gtctcatcta ccagtactgg tgtagcgtcc agtgaacaag 1980 acgcaacaga tcctgcttct gctaagccag tattcggcaa acccgcgttc ggagctattg 2040 ccaaagaacc gtcaacatca gaatatgcct ttggcaagcc atcttttggt gctccctcct 2100 ttggctctgg aaagtcatct gttgaatcgc ctgcctccgg atctgccttt ggtaagccct 2160 cttttggtac tccttccttt ggctctggaa attcatctgt tgagccgcct gcctccggat 2220 ctgcatttgg taagccctct tttggtactc cttcctttgg ctctggaaat tcatctgctg 2280 agccgcctgc ttccggatct gcctttggta agccctcttt tggtacatct gcattcggaa 2340 ctgcatcaag taacgaaact aactctggat ccatatttgg aaaggctgca tttggttcat 2400 catcttttgc accegecaac aatgaacttt teggateaaa etttaetatt teaaaaceta 2460

Printed:17-08-2000

SEUL

cagttgacag	cccaaaggag	gtagattcaa	cgtcaccttt	cccatcttct	ggcgatcaaa	2520
gtgaagatga	gtctaagagt	gatgtagact	cttcttcgac	accttttggt	acgaaaccta	2580
acacctctac	gaaaccaaag	accaatgcct	ttgattttgg	gagttcttcc	tttggatctg	2640
gattttcaaa	ggctctggaa	tctgttggtt	ccgatacaac	ttttaaattc	ggtactcagg	2700
cttcaccttt	ctcttcacag	ttaggaaaca	aatcaccatt	cagttccttc	acaaaagatg	2760
atactgaaaa	tggatcttta	agtaagggct	ctaccagtga	aatcaatgac	gataatgaag	2820
aacacgaaag	caatggtccc	aacgtaagcg	gtaatgattt	gacagattct	acggttgagc	2880
aaacatcttc	tactagatta	ccggaaactc	cctcggatga	agatggtgaa	gttgtcgagg	2940
aggaagcgca	aaaatccccc	ataggcaagc	taactgaaac	tataaaaaaa	agtgccaata	3000
ttgacatggc	tggtttaaaa	aatcctgtat	ttggaaatca	tgtcaaagca	aaatccgaat	3060
cgccgttttc	agcatttgca	acaaatatta	ccaaaccaag	ctctacaaca	cctgcttttt	3120
cgtttggtaa	ctccacaatg	aataaaagta	atacatctac	ggtttcacca	atggaagaag	3180
ctgatactaa	agaaactagt	gaaaagggcc	ccataacctt	gaagagtgtg	gagaatccgt	3240
ttctaccagc	gaaagaagaa	agaactggag	aaagttctaa	aaaggatcat	aacgatgacc	3300
caaaagatgg	ttatgtatca	ggaagtgaaa	tatctgtaag	gacttctgaa	agtgcttttg	3360
ataccacagc	aaacgaagaa	attccaaagt	cacaggacgt	gaacaatcat	gaaaaaagcg	3420
aaacagaccc	aaaatatagt	caacatgctg	tggttgatca	cgataacaag	tctaaagaaa	3480
tgaatgaaac	ttcgaagaat	aatgaaagga	gcggtcaacc	aaatcatggt	gtccaaggag	3540
atggaatagc	attgaaaaaa	gacaatgaaa	aagagaattt	tgattcaaat	atggcaataa	3600
agcaattcga	agaccaccaa	tcttcagaag	aggacgcgag	cgaaaaagac	agtagacaaa	3660
					gatgaaagta	
					catggtggag	
					acttcattag	
					aaggaaaatg	
					aagaaggaag	
					gttcaaacat	
					aaagaatatt	
					ttaaggttga	
					aacatccgga	
					acagtgcggt	
					aatattcaga	
					ctgaaggaaa	
					gccaatgcaa	
					gttaaagatc	
					gatgtgtccg	
	_			_	gtaaagaata	
					gcacgtgacg	
					ttggaggaga atgaaaggat	
		• •			gatttcttca	
	catggcaaaa		ucucyaaada	geadactyge	gattettea	4883
aaaacccyaa	carggradaa	9				1000

<210> 136

<211> 1460

<212> PRT

<213> Saccharomyces cerevisiae

<400> 136

Met Ser Ser Leu Lys Asp Glu Val Pro Thr Glu Thr Ser Glu Asp Phe
1 5 10 15

Gly Phe Lys Phe Leu Gly Gln Lys Gln Ile Leu Pro Ser Phe Asn Glu 20 25 30

Lys Leu Pro Phe Ala Ser Leu Gln Asn Leu Asp Ile Ser Asn Ser Lys 35 40 45

Ser Leu Phe Val Ala Ala Ser Gly Ser Lys Ala Val Val Gly Glu Leu 50 55 60

Gln Leu Leu Arg Asp His Ile Thr Ser Asp Ser Thr Pro Leu Thr Phe 65 70 75 80

Lys Trp Glu Lys Glu Ile Pro Asp Val Ile Phe Val Cys Phe His Gly 85 90 95

Asp Gln Val Leu Val Ser Thr Arg Asn Ala Leu Tyr Ser Leu Asp Leu 100 105 110

Glu Glu Leu Ser Glu Phe Arg Thr Val Thr Ser Phe Glu Lys Pro Val 115 120 125

Phe Gln Leu Lys Asn Val Asn Asn Thr Leu Val Ile Leu Asn Ser Val 130 135 140

Asn Asp Leu Ser Ala Leu Asp Leu Arg Thr Lys Ser Thr Lys Gln Leu 145 150 155 160

Ala Gln Asn Val Thr Ser Phe Asp Val Thr Asn Ser Gln Leu Ala Val 165 170 175

Leu Leu Lys Asp Arg Ser Phe Gln Ser Phe Ala Trp Arg Asn Gly Glu 180 185 190

Met Glu Lys Gln Phe Glu Phe Ser Leu Pro Ser Glu Leu Glu Glu Leu
195 200 205

Pro Val Glu Glu Tyr Ser Pro Leu Ser Val Thr Ile Leu Ser Pro Gln 210 215 220

Asp Phe Leu Ala Val Phe Gly Asn Val Ile Ser Glu Thr Asp Asp Glu 225 230 235 240

Val Ser Tyr Asp Gln Lys Met Tyr Ile Ile Lys His Ile Asp Gly Ser 245 250 255 Ala Ser Phe Gln Glu Thr Phe Asp Ile Thr Pro Pro Phe Gly Gln Ile 260 270 Val Arg Phe Pro Tyr Met Tyr Lys Val Thr Leu Ser Gly Leu Ile Glu 275 280 285 Pro Asp Ala Asn Val Asn Val Leu Ala Ser Ser Cys Ser Ser Glu Val 295 300 Ser Ile Trp Asp Ser Lys Gln Val Ile Glu Pro Ser Gln Asp Ser Glu 310 315 Arg Ala Val Leu Pro Ile Ser Glu Glu Thr Asp Lys Asp Thr Asn Pro 325 330 Ile Gly Val Ala Val Asp Val Val Thr Ser Gly Thr Ile Leu Glu Pro 345 Cys Ser Gly Val Asp Thr Ile Glu Arg Leu Pro Leu Val Tyr Ile Leu 355 360 365 Asn Asn Glu Gly Ser Leu Gln Ile Val Gly Leu Phe His Val Ala Ala 370 375 380 Ile Lys Ser Gly His Tyr Ser Ile Asn Leu Glu Ser Leu Glu His Glu 390 395 Lys Ser Leu Ser Pro Thr Ser Glu Lys Ile Pro Ile Ala Gly Gln Glu 410 Gln Glu Glu Lys Lys Lys Asn Asn Glu Ser Ser Lys Ala Leu Ser Glu 420 425 Asn Pro Phe Thr Ser Ala Asn Thr Ser Gly Phe Thr Phe Leu Lys Thr 435 440 445 Gln Pro Ala Ala Ala Asn Ser Leu Gln Ser Gln Ser Ser Ser Thr Phe 450 455

Gly Ala Pro Ser Phe Gly Ser Ser Ala Phe Lys Ile Asp Leu Pro Ser 465 470 475 480

Val Ser Ser Thr Ser Thr Gly Val Ala Ser Ser Glu Gln Asp Ala Thr
485 490 495

Asp Pro Ala Ser Ala Lys Pro Val Phe Gly Lys Pro Ala Phe Gly Ala 500 505 510

Ile	Ala	Lys 515	Glu	Pro	Ser	Thr	Ser 520	Glu	Tyr	Ala	Phe	Gly 525	Lys	Pro	Ser	
Phe	Gly 530	Ala	Pro	Ser	Phe	Gly 535	Ser	Gly	Lys	Ser	Ser 540	Val	Glu	Ser	Pro	
Ala 545	Ser	Gly	Ser	Ala	Phe 550	Gly	Lys	Pro	Ser	Phe 555	Gly	Thr	Pro	Ser	Phe 560	
Gly	Ser	Gly	Asn	Ser 565	Ser	Val	Glu	Pro	Pro 570	Ala	Ser	Gly	Ser	Ala 575	Phe	
Gly	Lys	Pro	Ser 580	Phe	Gly	Thr	Pro	Ser 585	Phe	Gly	Ser	Gly	Asn 590	Ser	Ser	
		595				_	600					605		Phe		
	610					615					620			Gly		
625			_		630					635				Ala	640	
				645					650					Val 655		
			660					665					670	Gly		
		675	_				680					685		Thr		
	690					695					700			Ala		
705					710					715				Ser	720	
				725					730					735		
			740					745					750	Thr		
ASP	ASP	Thr	GIU	ASN	стХ	ser	Leu 260		пÃg	σтλ	Ser	765	261	Glu	T16	

760

Asn Asp Asp Asn Glu Glu His Glu Ser Asn Gly Pro Asn Val Ser Gly Asn Asp Leu Thr Asp Ser Thr Val Glu Gln Thr Ser Ser Thr Arg Leu 795 -Pro Glu Thr Pro Ser Asp Glu Asp Gly Glu Val Val Glu Glu Glu Ala Gln Lys Ser Pro Ile Gly Lys Leu Thr Glu Thr Ile Lys Lys Ser Ala Asn Ile Asp Met Ala Gly Leu Lys Asn Pro Val Phe Gly Asn His Val Lys Ala Lys Ser Glu Ser Pro Phe Ser Ala Phe Ala Thr Asn Ile Thr Lys Pro Ser Ser Thr Thr Pro Ala Phe Ser Phe Gly Asn Ser Thr Met Asn Lys Ser Asn Thr Ser Thr Val Ser Pro Met Glu Glu Ala Asp Thr Lys Glu Thr Ser Glu Lys Gly Pro Ile Thr Leu Lys Ser Val Glu Asn Pro Phe Leu Pro Ala Lys Glu Glu Arg Thr Gly Glu Ser Ser Lys Lys Asp His Asn Asp Asp Pro Lys Asp Gly Tyr Val Ser Gly Ser Glu Ile Ser Val Arg Thr Ser Glu Ser Ala Phe Asp Thr Thr Ala Asn Glu Glu Ile Pro Lys Ser Gln Asp Val Asn Asn His Glu Lys Ser Glu Thr Asp Pro Lys Tyr Ser Gln His Ala Val Val Asp His Asp Asn Lys Ser Lys Glu Met Asn Glu Thr Ser Lys Asn Asn Glu Arg Ser Gly Gln Pro Asn

His Gly Val Gln Gly Asp Gly Ile Ala Leu Lys Lys Asp Asn Glu Lys 

- Glu Asn Phe Asp Ser Asn Met Ala Ile Lys Gln Phe Glu Asp His Gln 1025 1030 1035 1040
- Ser Ser Glu Glu Asp Ala Ser Glu Lys Asp Ser Arg Gln Ser Ser Glu
  1045 1050 1055
- Val Lys Glu Ser Asp Asp Asn Met Ser Leu Asn Ser Asp Arg Asp Glu 1060 1065 1070
- Ser Ile Ser Glu Ser Tyr Asp Lys Leu Glu Asp Ile Asn Thr Asp Glu 1075 1080 1085
- Leu Pro His Gly Gly Glu Ala Phe Lys Ala Arg Glu Val Ser Ala Ser 1090 1095 1100
- Ala Asp Phe Asp Val Gln Thr Ser Leu Glu Asp Asn Tyr Ala Glu Ser 1105 1110 1115 1120
- Gly Ile Gln Thr Asp Leu Ser Glu Ser Ser Lys Glu Asn Glu Val Gln 1125 1130 1135
- Thr Asp Ala Ile Pro Val Lys His Asn Ser Thr Gln Thr Val Lys Lys 1140 1145 1150
- Glu Ala Val Asp Asn Gly Leu Gln Thr Glu Pro Val Glu Thr Cys Asn 1155 1160 1165
- Phe Ser Val Gln Thr Phe Glu Gly Asp Glu Asn Tyr Leu Ala Glu Gln 1170 1175 1180
- Cys Lys Pro Lys Gln Leu Lys Glu Tyr Tyr Thr Ser Ala Lys Val Ser 1185 1190 1195 1200
- Asn Ile Pro Phe Val Ser Gln Asn Ser Thr Leu Arg Leu Ile Glu Ser 1205 1210 1215
- Thr Phe Gln Thr Val Glu Ala Glu Phe Thr Val Leu Met Glu Asn Ile 1220 1225 1230
- Arg Asn Met Asp Thr Phe Phe Thr Asp Gln Ser Ser Ile Pro Leu Val 1235 1240 1245
- Lys Arg Thr Val Arg Ser Ile Asn Asn Leu Tyr Thr Trp Arg Ile Pro 1250 1260
- Glu Ala Glu Ile Leu Leu Asn Ile Gln Asn Asn Ile Lys Cys Glu Gln 1265 1270 1275 1280

Met Gln Ile Thr Asn Ala Asn Ile Gln Asp Leu Lys Glu Lys Val Thr 1285 1290 1295

Asp Tyr Val Arg Lys Asp Ile Ala Gln Ile Thr Glu Asp Val Ala Asn 1300 1305 1310

Ala Lys Glu Glu Tyr Leu Phe Leu Met His Phe Asp Asp Ala Ser Ser 1315 1320 1325

Gly Tyr Val Lys Asp Leu Ser Thr His Gln Phe Arg Met Gln Lys Thr
1330 1335 1340

Leu Arg Gln Lys Leu Phe Asp Val Ser Ala Lys Ile Asn His Thr Glu 1345 1350 1355 1360

Glu Leu Leu Asn Ile Leu Lys Leu Phe Thr Val Lys Asn Lys Arg Leu 1365 1370 1375

Asp Asp Asn Pro Leu Val Ala Lys Leu Ala Lys Glu Ser Leu Ala Arg 1380 1385 1390

Asp Gly Leu Leu Lys Glu Ile Lys Leu Leu Arg Glu Gln Val Ser Arg 1395 1400 1405

Leu Gln Leu Glu Glu Lys Gly Lys Lys Ala Ser Ser Phe Asp Ala Ser 1410 1415 1420

Ser Ser Ile Thr Lys Asp Met Lys Gly Phe Lys Val Val Glu Val Gly 1425 1430 1435 1440

Leu Ala Met Asn Thr Lys Lys Gln Ile Gly Asp Phe Phe Lys Asn Leu 1445 1450 1455

Asn Met Ala Lys 1460

<210> 137

<211> 1321

<212> DNA

<213> Saccharomyces cerevisiae

<400> 137

tcgatcaact ctatccaaca attctataat atccactgtt cattaacgaa tattggtctt 60 tttcccttat ggtgaagtaa attttccatg caatatccgg gtaagctatc gacaagttta 120 ttgactgcaa tttgagttta ttacatccgt acattactaa gatgtatggt ttttctattt 180

```
ttgtgctccg tgtcgatatt tcgtggagca aaccagaaaa gatgcggaac ctcttagcac 240
tccgcctgga cataggcgga gcatattcct cctatgggat gggttttgtt gtactctttt 300
ctctctagac aggacctccg attgcctccc tgagggtgag atggtttccg gcctcaggac 360
ggccttctcc agtttctagc gaggcataca ttccaaccaa aggtgtatca agaatatctg 420
aaattaaagg tagttgaatc tctatttgtt gttgttatta ccgcttatta tcccatagtt 480
gagacgacca agattcaaac atgcaaatgt atgcaccata tccattctaa acatagtttt 540
tcgaacgttc agagcttaaa gggacaatta ttttagaaac tgaattttta cccagtggaa 600
taacatcgta tctgtaaagt ctacaaaatt ttttatccat caaaaattaa aacaaagaaa 660
actgccaaac tgaatatgag gaactttcct ctctaggaat gacttagtga atgtacagtg 720
acttgtggaa aatatgatta gattttgagc gggtgatgcg acttaacagt ctcattgcct 780
aagaaatatc caaatttgtg gttcatgctc tcccccaaga tatgacgatg agagctcgtt 840
taaaattttg tcttcttccg aacagttatg aaaaaactat tacgtgtttt tatgatatcc 900
ttactaactt gtcatttttt tataaaatta tttttttaac agttttgtca agactttgac 960
tggtaagacc atcactttgg aagttgaatc ttctgacact attgacaatg tcaagtcaaa 1020
gattcaagac aaggaaggta tcccacctga ccaacaaaga ttgatctttg ctggtaagca 1080
attggaagac ggtagaacct tgtctgacta caacattcaa aaagaatcca ctttgcactt 1140
agtettgaga ttgagaggtg gtateattga accatetttg aaagetttgg ettecaagta 1200
caactgtgac aaatctgttt gccgtaagtg ttatgctaga ttgccaccaa gagctaccaa 1260
ctgtagaaag agaaagtgtg gtcacaccaa ccaattgcgt ccaaagaaga agttaaaatg 1320
                                                                  1321
```

```
<210> 138
```

<211> 128 ----

<212> PRT

<213> Saccharomyces cerevisiae

<400> 138

Met Gln Ile Phe Val Lys Thr Leu Thr Gly Lys Thr Ile Thr Leu Glu
1 5 10 15

Val Glu Ser Ser Asp Thr Ile Asp Asn Val Lys Ser Lys Ile Gln Asp
20 25 30

Lys Glu Gly Ile Pro Pro Asp Gln Gln Arg Leu Ile Phe Ala Gly Lys
35 40 45

Gln Leu Glu Asp Gly Arg Thr Leu Ser Asp Tyr Asn Ile Gln Lys Glu
50 55 60

Ser Thr Leu His Leu Val Leu Arg Leu Arg Gly Gly Ile Ile Glu Pro 65 70 75 80

Ser Leu Lys Ala Leu Ala Ser Lys Tyr Asn Cys Asp Lys Ser Val Cys 85 90 95

Arg Lys Cys Tyr Ala Arg Leu Pro Pro Arg Ala Thr Asn Cys Arg Lys
100 105 110



Arg Lys Cys Gly His Thr Asn Gln Leu Arg Pro Lys Lys Leu Lys 115 120 125

<210> 139 <211> 2216 <212> DNA <213> Saccharomyces cerevisiae

<400> 139

aatattcaac ctgaagttgc agtcaaagcg gcgattaact tttccaacgt aaccgatcta 60 actaacaata gcaccgatgg agcaaaaata acagaaattg gaagtacatc taaacggcca 120 attgagagcg gtacatcttc tgatccagac accaaaaagg ttaaagagag tccagcaaat 180 gatcaagctt ccaacgagtg atgtaatatt aaacaatgta attatataaa tatgaaacat 240 ctacatattt taaatgtcac taatgtcatt acagaggaca taaagtgatt tatgacacat 300 ccgtactagt agttaagtat gaacaaattt tgggtttatt tgccattttt tttcacgcgg 360 gtttcttgga tgcgcaaacc caccttttct aacaccacta agaaatatca actttatagg 420 ccatcgaaga taaaggaacg taagtttgtc aattcaacct cacattttca acgcacatta 480 agcacttggt tcgtggagaa atgaatgatc ctcgtgaaat tttagcggtt gatccgtaca 540 ataatattac ttctgatgaa gaggatgagc aagccatcgc gagagaactt gaatttatgg 600 aacgaaagag gcaggcctta gtggaacgat taaaaagaaa gcaagaattt aagaaacccc 660 aggatectaa ttttgaagee ategaggtae etcaatetee taccaaaaae egtgtgaaag 720 tggggtctca taatgctaca caacaaggca caaaattcga aggttcgaat attaatgaag 780 taaggttatc tcaattacag cagcaaccaa aaccaccagc tagtacaacc acatacttta 840 tggagaaatt tcaaaacgca aagaagaacg aagataaaca aattgccaag tttgaaagca 900 tgatgaatgc aagagtacat acgttcagta ccgatgagaa gaaatatgtg ccgataatca 960 caaacgaatt agaaagcttt tcaaatcttt gggttaaaaa gaggtacata cctgaagatg 1020 acttaaaacg ggctttgcat gagatcaaaa tccttcggtt gggcaaactt tttgctaaaa 1080 ttcgcccacc taaatttcaa gagcctgaat acgccaactg ggccaccgta ggcctcatta 1140 gccacaaatc ggacatcaaa tttacatcat ctgaaaagcc agtcaaattc ttcatgttca 1200 ccataacgga ctttcagcat acactagatg tttatatctt cgggaaaaaag ggtgtagaaa 1260 gatattataa tettegeetg ggtgatgtga tagcaatatt aaacccagaa gtactaccat 1320 ggagaccete agggegagga aattttatea aateetteaa eettegaatt agteatgaet 1380 tcaaatgtat cctggagata ggttcaagta gagatttagg ttggtgtccc atagtgaata 1440 aaaagactca caaaaaatgt ggctctccca ttaacatatc tcttcataag tgttgcgatt 1500 accatagaga agtgcaattt cgtggaacaa gtgctaaaag aattgaatta aatggtgggt 1560 acgccttggg cgcgcctacg aaagtggact ctcaaccaag cctatataag gccaaagggg 1620 aaaacgggtt taatataatc aaaggtactc gtaagcgcct gtcagaagag gaggaaagac 1680 ttaaaaagag ctctcacaat tttacgaata gtaattctgc caaagcattt ttcgacgaga 1740 aatttcagaa tccagatatg ctggcaaact tagacaataa aagaaggaaa ataatagaaa 1800 ctaagaaatc gacagcactg agccgcgaac taggcaaaat tatgagaagg agggaatcca 1860 gcggattaga agataagagc gtcggagagc gacagaaaat gaaacgaacc acagaaagtg 1920 ccctccagac agggcttatc caacgcctag gattcgatcc cactcatgga aaaatttccc 1980

agaacgaatg gttcaagaaa agaagccatc gcgaagaagt ttggcaaaaa catttcggat 2160

ccaaggaaac taaagaaact tctgacggta gtgccagcga tcttgagata atataa

aagtactcaa gtcttctgta tcagggagcg aacctaagaa caacttactc ggtaaaaaaa 2040 aaactgttat aaacgatctc ttgcattaca agaaggaaaa agtcattctc gcaccttcaa 2100

<210> 140

<211> 571

<212> PRT

<213> Saccharomyces cerevisiae

<400> 140

Met Asn Asp Pro Arg Glu Ile Leu Ala Val Asp Pro Tyr Asn Asn Ile 1 5 10 15

Thr Ser Asp Glu Glu Asp Glu Gln Ala Ile Ala Arg Glu Leu Glu Phe
20 25 30

Met Glu Arg Lys Arg Gln Ala Leu Val Glu Arg Leu Lys Arg Lys Gln 35. 40 45

Glu Phe Lys Lys Pro Gln Asp Pro Asn Phe Glu Ala Ile Glu Val Pro 50 55 60

Gln Ser Pro Thr Lys Asn Arg Val Lys Val Gly Ser His Asn Ala Thr
65 70 75 80

Gln Gln Gly Thr Lys Phe Glu Gly Ser Asn Ile Asn Glu Val Arg Leu 85 90 95

Ser Gln Leu Gln Gln Gln Pro Lys Pro Pro Ala Ser Thr Thr Tyr
100 105 110

Phe Met Glu Lys Phe Gln Asn Ala Lys Lys Asn Glu Asp Lys Gln Ile 115 120 125

Ala Lys Phe Glu Ser Met Met Asn Ala Arg Val His Thr Phe Ser Thr 130 135 140

Asp Glu Lys Lys Tyr Val Pro Ile Ile Thr Asn Glu Leu Glu Ser Phe 145 150 155 160

Ser Asn Leu Trp Val Lys Lys Arg Tyr Ile Pro Glu Asp Asp Leu Lys 165 170 175

Arg Ala Leu His Glu Ile Lys Ile Leu Arg Leu Gly Lys Leu Phe Ala 180 185 190

												•	2		
Lys	Ile	Arg 195	Pro	Pro	Lys	Phe	Gln 200	Glu	Pro	Glu	Tyr	Ala 205	Asn	Trp	Ala
Thr	Val 210	Gly	Leu	Ile	Ser	His 215	Lys	Ser	Asp	Ile	Lys 220	Phe	Thr	Ser	Ser
Glu 225	Lys	Pro	Val	Lys	Phe 230	Phe	Met	Phe	Thr	Ile 235	Thr	Asp	Phe	Gln	His 240
Thr	Leu	Asp	Val	Tyr 245	Ile	Phe	Gly	Lys	Lys 250	Gly	Val	Glu	Arg	Tyr 255	туг
Asn	Leu	Arg	Leu 260	Gly	Asp	Val	Ile	Ala 265	Ile	Leu	Asn	Pro	Glu 270	Val	Leu
Pro	Trp	Arg 275	Pro	Ser	Gly	Arg	Gly 280	Asn	Phe	Ile	Lys	Ser 285	Phe	Asn	Leu
Arg	Ile 290	Ser	His	Asp	Phe	Lys 295	Cys	Ile	Leu	Glu	Ile 300	Gly	Ser	Ser	Arg
Asp 305	Leu	Gly	Trp	Cys	Pro 310	Ile	Val	Asn	Lys	Lys 315	Thr	His	Lys	Lys	Cys 320
Gly	Ser	Pro	Ile	Asn 325	Ile	Ser	Leu	His	Lys 330	Cys	Cys	Asp	Tyr	His 335	Arg
Glu	Val	Gln	Phe 340	Arg	Gly	Thr	Ser	Ala 345	Lys	Arg	Ile	Glu	Leu 350	Asn	Gly
Gly	Tyr	Ala 355	Leu	Gly	Ala	Pro	Thr 360	Lys	Val	Asp	Ser	Gln 365	Pro	Ser	Leu
Tyr	Lys 370	Ala	Lys	Gly	Glu	Asn 375	Gly	Phe	Asn	Ile	Ile 380	Lys	Gly	Thr	Arg
Lys 385	Arg	Leu	Ser	Glu	Glu 390	Glu	Glu	Arg	Leu	Lys 395	Lys	Ser	Ser	His	Asn 400
Phe	Thr	Asn	Ser		Ser		_				_		_	Phe	

Asn Pro Asp Met Leu Ala Asn Leu Asp Asn Lys Arg Arg Lys Ile Ile

Glu Thr Lys Lys Ser Thr Ala Leu Ser Arg Glu Leu Gly Lys Ile Met

```
Arg Arg Arg Glu Ser S r Gly Leu Glu Asp Lys Ser Val Gly Glu Arg
                                             460
    45.0
                        455
Gln Lys Met Lys Arg Thr Thr Glu Ser Ala Leu Gln Thr Gly Leu Ile
465
                    470
                                         475
Gln Arg Leu Gly Phe Asp Pro Thr His Gly Lys Ile Ser Gln Val Leu
                                     490
                485
Lys Ser Ser Val Ser Gly Ser Glu Pro Lys Asn Asn Leu Leu Gly Lys
                                 505
Lys Lys Thr Val Ile Asn Asp Leu Leu His Tyr Lys Lys Glu Lys Val
                            520
Ile Leu Ala Pro Ser Lys Asn Glu Trp Phe Lys Lys Arg Ser His Arg
    530
                        535
                                             540
Glu Glu Val Trp Gln Lys His Phe Gly Ser Lys Glu Thr Lys Glu Thr
545
                    550
                                                              560
Ser Asp Gly Ser Ala Ser Asp Leu Glu Ile Ile
                565
                                     570
<210> 141
<211> 1133
<212> DNA
<213> Saccharomyces cerevisiae
```

<400> 141

```
gacattttga aaaaattaca tttatagaag tttattataa cgtaacagtt aacaacaaag 60
gctcatagtc cgagatcaat ataataaatt attcaaggtt taaggaagaa gttaccatgg 120
aaatgactta ctatgaaaag acacctttga ttcgtcaatt tttgaacaat ggtaagacaa 180
attcgtggtt ttacgttaag catgagatgt tacaaccagg tggaagtttc aaatcgagag 240
gaatcgggca tttgataagg aagagtaatg aagaagcgct aagcgagggt tctgggaagc 300
ttgctgtatt ttctagctct gggggaaatg ctggtttagc agcagcaact gcctgcagat 360
cgatggcact taattgcagt gtagtggttc ctaaaactac aaaacctaga atggtaaaga 420
aaattcaaag tgcaggagcc aaagtcatta tccatggtga tcattggggg gaagcagatg 480
aatacttgag gcacgaatga atggcgcaag aaagccaaca tggttcgaag acactatatg 540
tgcacccgtt tgataacgag acaatttggg aaggtcattc tacgattgtg gatgaaatca 600
tagaacaatt gaaggaaaat gatatatcct tacctagggt gaaagctttg gtttgtagtg 660
ttggtggtgg tgggctattt agtggcataa ttaaaggcct agataggaat cagcttgctg 720
aaaaaattcc ggtcgttgct gtagaaactg ccggttgtga cgtattgaat aagtctctca 780
aaaaaggtag tccagttact cttgaaaaat tgacaagtgt tgcaacttct ttggcctccc 840
catacatagc atcattcgcg tttgagagtt ttaacaagta tggatgtaag tctgtagttt 900
tatcagatca agacgttctg gcaacatgct tgagatatgc cgatgactac aattttatag 960
```

tggaaccagc ctgtggagca tccttacatt tatgttatca tccagagatt cttgaagaca 1020 ttctggaaca aaaaatatat gaggatgata tcgttattat aatcgcatgc ggtggatcat 1080 gtatgacgta tgaagacttg gtgaaagcgt cgagcacatt aaacgtatca taa 1133

<210> 142

<211> 210

<212> PRT

<213> Saccharomyces cerevisiae

<400> 142

Met Ala Gln Glu Ser Gln His Gly Ser Lys Thr Leu Tyr Val His Pro 1 5 10 15

Phe Asp Asn Glu Thr Ile Trp Glu Gly His Ser Thr Ile Val Asp Glu 20 25 30

Ile Ile Glu Gln Leu Lys Glu Asn Asp Ile Ser Leu Pro Arg Val Lys
35 40 45

Ala Leu Val Cys Ser Val Gly Gly Gly Leu Phe Ser Gly Ile Ile
50 55 60

Lys Gly Leu Asp Arg Asn Gln Leu Ala Glu Lys Ile Pro Val Val Ala 65 70 75 80

Val Glu Thr Ala Gly Cys Asp Val Leu Asn Lys Ser Leu Lys Lys Gly 85 90 95

Ser Pro Val Thr Leu Glu Lys Leu Thr Ser Val Ala Thr Ser Leu Ala 100 105 110

Ser Pro Tyr Ile Ala Ser Phe Ala Phe Glu Ser Phe Asn Lys Tyr Gly
115 120 125

Cys Lys Ser Val Val Leu Ser Asp Gln Asp Val Leu Ala Thr Cys Leu 130 135 140

Arg Tyr Ala Asp Asp Tyr Asn Phe Ile Val Glu Pro Ala Cys Gly Ala 145 150 155 160

Ser Leu His Leu Cys Tyr His Pro Glu Ile Leu Glu Asp Ile Leu Glu
165 170 175

Gln Lys Ile Tyr Glu Asp Asp Ile Val Ile Ile Ile Ala Cys Gly Gly
180 185 190

Ser Cys Met Thr Tyr Glu Asp Leu Val Lys Ala Ser Ser Thr Leu Asn

SEQL\*

195 200 205

Val Ser 210

<210> 143 <211> 2549 <212> DNA <213> Saccharomyces cerevisiae

<400> 143

ccatgaactc agcatgtgct actccagtta atgacttgtt cgtatcgttc atgccataag 60 ccatcacctg gccagttggc gtatgtacaa agatgcaagc taccggtgtc tcatcgtggt 120 caagagcgta tctagccaaa cggacagctg tcctcatatg tttaatatgc tgcatagtgt 180 gagtcctcta gtttttaccg cagccaccag ccgcttctcg agcaaagtgt agatcccatt 240 aggactcatc attcatctaa ttttgctatg ttagctgcaa ctttctattt taatagaacc 300 ttctggaaat ttcacccggc gcggcacccg aggaactgga cagcgtgtcg aaaaagttgc 360 ttttttatat aaaggacacg aaaagggttc tctggaagat ataaatatgg ctatgtaatt 420 ctaaagatta acgtgttact gttttacttt tttaaagtcc ccaagagtag tctcaaggga 480 aaaagcgtat caaacatacc atgtttttca acagactaag cgctggcaag ctgctggtac 540 cactctccgt ggtcctgtac gcccttttcg tggtaatatt acctttacag aattctttcc 600 actcctccaa tgttttagtt agaggtgccg atgatgtaga aaactacgga actgttatcg 660 gtattgactt aggtactact tattcctgtg ttgctgtgat gaaaaatggt aagactgaaa 720 ttcttgctaa tgagcaaggt aacagaatca ccccatctta cgtggcattc accgatgatg 780 aaagattgat tggtgatgct gcaaagaacc aagttgctgc caatcctcaa aacaccatct 840 tcgacattaa gagattgatc ggtttgaaat ataacgacag atctgttcag aaggatatca 900 agcacttgcc atttaatgtg gttaataaag atgggaagcc cgctgtagaa gtaagtgtca 960 aaggagaaaa gaaggttttt actccagaag aaatttctgg tatgatcttg ggtaagatga 1020 aacaaattgc cgaagattat ttaggcacta aggttaccca tgctgtcgtt actgttcctg 1080 cttatttcaa tgacgcgcaa agacaagcca ccaaggatgc tggtaccatc gctggtttga 1140 acgttttgag aattgttaat gaaccaaccg cagccgccat tgcctacggt ttggataaat 1200 ctgataagga acatcaaatt attgtttatg atttgggtgg tggtactttc gatgtctctc 1260 tattgtctat tgaaaacggt gttttcgaag tccaagccac ttctggtgat actcatttag 1320 gtggtgaaga ttttgactat aagatcgttc gtcaattgat aaaagctttc aagaagaagc 1380 atggtattga tgtgtctgac aacaacaagg ccctagctaa attgaagaga gaagctgaaa 1440 aggetaaacg tgccttgtcc agccaaatgt ccacccgtat tgaaattgac tccttcgttg 1500 atggtatcga cttaagtgaa accttgacca gagctaagtt tgaggaatta aacctagatc 1560 tattcaagaa gaccttgaag cctgtcgaga aggttttgca agattctggt ttggaaaaga 1620 aggatgttga tgatatcgtt ttggttggtg gttctactag aattccaaag gtccaacaat 1680 tgttagaatc atactttgat ggtaagaagg cctccaaggg tattaaccca gatgaagctg 1740 ttgcatacgg tgcagccgtt caagctggtg tcttatccgg tgaagaaggt gtcgaagata 1800 ttgttttatt ggatgtcaac gctttgactc ttggtattga aaccactggt ggtgtcatga 1860 ctccattaat taagagaaat actgctattc ctacaaagaa atcccaaatt ttctctactg 1920 ccgttgacaa ccaaccaacc gttatgatca aggtatacga gggtgaaaga gccatgtcta 1980 aggacaacaa tctattaggt aagtttgaat taaccggcat tccaccagca ccaagaggtg 2040 tacctcaaat tgaagtcaca tttgcacttg acgctaatgg tattctgaag gtgtctgcca 2100

cagataaggg aactggtaaa tccgaatcta tcaccatcac taacgataaa ggtagattaa 2160 cccaagaaga gattgataga atggttgaag aggctgaaaa attcgcttct gaagacgctt 2220 ctatcaaggc caaggttgaa tctagaaaca aattagaaaa ctacgctcac tctttgaaaa 2280 accaagttaa tggtgaccta ggtgaaaaat tggaagaaga agacaaggaa accttattag 2340 atgctgctaa cgatgttta gaatggttag atgataactt tgaaaccgcc attgctgaag 2400 actttgatga aaagttcgaa tctttgtcca aggtcgctta tccaattact tctaagttgt 2460 acggaggtgc tgatggttct ggtgccgctg attatgacga cgaagatgaa gatgacgatg 2520 gtgattattt cgaacacgac gaattgtag

<210> 144

<211> 682

<212> PRT

<213> Saccharomyces cerevisiae

<400> 144

Met Phe Phe Asn Arg Leu Ser Ala Gly Lys Leu Leu Val Pro Leu Ser 1 5 10 15

Val Val Leu Tyr Ala Leu Phe Val Val Ile Leu Pro Leu Gln Asn Ser 20 25 30

Phe His Ser Ser Asn Val Leu Val Arg Gly Ala Asp Asp Val Glu Asn 35 40 45

Tyr Gly Thr Val Ile Gly Ile Asp Leu Gly Thr Thr Tyr Ser Cys Val 50 55 60

Ala Val Met Lys Asn Gly Lys Thr Glu Ile Leu Ala Asn Glu Gln Gly 65 70 75 80

Asn Arg Ile Thr Pro Ser Tyr Val Ala Phe Thr Asp Asp Glu Arg Leu 85 90 95

Ile Gly Asp Ala Ala Lys Asn Gln Val Ala Ala Asn Pro Gln Asn Thr
100 105 110

Ile Phe Asp Ile Lys Arg Leu Ile Gly Leu Lys Tyr Asn Asp Arg Ser 115 120 125

Val Gln Lys Asp Ile Lys His Leu Pro Phe Asn Val Val Asn Lys Asp 130 135 140

Gly Lys Pro Ala Val Glu Val Ser Val Lys Gly Glu Lys Lys Val Phe 145 150 155 160

Thr Pro Glu Glu Ile Ser Gly Met Ile Leu Gly Lys Met Lys Gln Ile 165 170 175

31a	Clu	λen	ጥህጉ	Leu	Glv	Thr	Lys	Val	Thr	His	Ala	Val	Val	Thr	Val
мта	Gru	ಬಾಗಿ	180					185					190		

- Pro Ala Tyr Phe Asn Asp Ala Gln Arg Gln Ala Thr Lys Asp Ala Gly
  195 200 205
- Thr Ile Ala Gly Leu Asn Val Leu Arg Ile Val Asn Glu Pro Thr Ala 210 215 220
- Ala Ala Ile Ala Tyr Gly Leu Asp Lys Ser Asp Lys Glu His Gln Ile 225 230 235 240
- Ile Val Tyr Asp Leu Gly Gly Gly Thr Phe Asp Val Ser Leu Leu Ser 245 250 255
- Ile Glu Asn Gly Val Phe Glu Val Gln Ala Thr Ser Gly Asp Thr His 260 265 270
- Leu Gly Gly Glu Asp Phe Asp Tyr Lys Ile Val Arg Gln Leu Ile Lys 275 280 285
- Ala Phe Lys Lys Lys His Gly Ile Asp Val Ser Asp Asn Asn Lys Ala 290 295 300
- Leu Ala Lys Leu Lys Arg Glu Ala Glu Lys Ala Lys Arg Ala Leu Ser 305 310 315 320
- Ser Gln Met Ser Thr Arg Ile Glu Ile Asp Ser Phe Val Asp Gly Ile 325 330 335
- Asp Leu Ser Glu Thr Leu Thr Arg Ala Lys Phe Glu Glu Leu Asn Leu 340 345 350
- Asp Leu Phe Lys Lys Thr Leu Lys Pro Val Glu Lys Val Leu Gln Asp 355 360 365
- Ser Gly Leu Glu Lys Lys Asp Val Asp Asp Ile Val Leu Val Gly Gly 370 375 380
- Ser Thr Arg Ile Pro Lys Val Gln Gln Leu Leu Glu Ser Tyr Phe Asp 385 390 395 400
- Gly Lys Lys Ala Ser Lys Gly Ile Asn Pro Asp Glu Ala Val Ala Tyr 405 410 415
- Gly Ala Ala Val Gln Ala Gly Val Leu Ser Gly Glu Glu Gly Val Glu 425 430

Asp Ile Val Leu Leu Asp Val Asn Ala Leu Thr Leu Gly Ile Glu Thr 435 440 445

Thr Gly Gly Val Met Thr Pro Leu Ile Lys Arg Asn Thr Ala Ile Pro 450 455 460

Thr Lys Lys Ser Gln Ile Phe Ser Thr Ala Val Asp Asn Gln Pro Thr 465 470 475 480

Val Met Ile Lys Val Tyr Glu Gly Glu Arg Ala Met Ser Lys Asp Asn 485 490 495

Asn Leu Leu Gly Lys Phe Glu Leu Thr Gly Ile Pro Pro Ala Pro Arg
500 505 510

Gly Val Pro Gln Ile Glu Val Thr Phe Ala Leu Asp Ala Asn Gly Ile 515 520 525

Leu Lys Val Ser Ala Thr Asp Lys Gly Thr Gly Lys Ser Glu Ser Ile 530 535 540

Thr Ile Thr Asn Asp Lys Gly Arg Leu Thr Gln Glu Glu Ile Asp Arg
545 550 555 560

Met Val Glu Glu Ala Glu Lys Phe Ala Ser Glu Asp Ala Ser Ile Lys 565 570 575

Ala Lys Val Glu Ser Arg Asn Lys Leu Glu Asn Tyr Ala His Ser Leu
580 585 590

Lys Asn Gln Val Asn Gly Asp Leu Gly Glu Lys Leu Glu Glu Glu Asp 595 600 605

Lys Glu Thr Leu Leu Asp Ala Ala Asn Asp Val Leu Glu Trp Leu Asp 610 620

Asp Asn Phe Glu Thr Ala Ile Ala Glu Asp Phe Asp Glu Lys Phe Glu 625 630 635 640

Ser Leu Ser Lys Val Ala Tyr Pro Ile Thr Ser Lys Leu Tyr Gly Gly 645 650 655

Ala Asp Gly Ser Gly Ala Ala Asp Tyr Asp Asp Glu Asp Glu Asp Asp 660 665 670

Asp Gly Asp Tyr Phe Glu His Asp Glu Leu 675 680

```
<210> 145
<211> 1253
<212> DNA
<213> Saccharomyces cerevisiae
<400> 145
tacctaagtc aataccgata acagttccgt agttttctac atcatcggca cctctaacta 60
aaacattgga ggagtggaaa gaattctgta aaggtaatat taccacgaaa agggcgtaca 120
ggaccacgga gagtggtacc agcagcttgc cagcgcttag tctgttgaaa aacatggtat 180
gtttgatacg ctttttccct tgagactact cttggggact ttaaaaaaagt aaaacagtaa 240
cacgitaatc titagaatta catagccata titatatcti ccagagaacc cititcgtgt 300
cetttatata aaaaagcaac tttttegaca egetgteeag tteetegggt geegegeegg 360
gtgaaatttc cagaaggttc tattaaaata gaaagttgca gctaacatag caaaattaga 420
tgaatgatga gtcctaatgg gatctacact ttgctcgaga agcggctggt ggctgcggta 480
aaaactagag gactcacact atgcagcata ttaaacatat gaggacagct gtccgtttgg 540
ctagatacgc tettgaccac gatgagacac eggtagettg catetttgta catacgccaa 600
ctggccaggt gatggcttat ggcatgaacg atacgaacaa gtcattaact ggagtagcac 660
atgctgagtt catggggatc gatcagatca aggcgatgtt gggctcccga ggagttgttg 720
acgtgttcaa agacattact ctatatgtta ctgtagaacc gtgtataatg tgtgcatctg 780
ctctcaagca attagacatt ggaaaggtgg tgttcggttg tggcaacgag agatttggag 840
gcaacggtac tgtcttgtca gtaaatcatg atacgtgtac attagtgccc aagaacaata 900
gtgcggcagg gtacgagagt ataccgggga tcttgaggaa agaagcaata atgctgctga 960
gatactttta tgtaagacaa aatgaaaggg cgccaaagcc acggtccaag agtgacagag 1020
tgttggataa aaacacgttt ccgcctatgg aatggtcaaa gtatcttaat gaagaagcat 1080
tcattgagac ttttggtgat gattacagga cttgttttgc gaataaagtt gacttgtcca 1140
gtaatagcgt cgattgggat ttgattgact cccaccaaga taatataatc caagaactgg 1200
                                                                  1253
aagaacaatg caaaatgttt aagtttaatg tacataagaa atctaaggtt tga
<210> 146
<211> 250
<212> PRT
<213> Saccharomyces cerevisiae
<400> 146
Met Gln His Ile Lys His Met Arg Thr Ala Val Arg Leu Ala Arg Tyr
  1
                                     10
Ala Leu Asp His Asp Glu Thr Pro Val Ala Cys Ile Phe Val His Thr
                                 25
             20
Pro Thr Gly Gln Val Met Ala Tyr Gly Met Asn Asp Thr Asn Lys Ser
                             40
```

Leu Thr Gly Val Ala His Ala Glu Phe Met Gly Ile Asp Gln Ile Lys

55

60

Ala Met Leu Gly Ser Arg Gly Val Val Asp Val Phe Lys Asp Ile Thr 65 70 75 80

Leu Tyr Val Thr Val Glu Pro Cys Ile Met Cys Ala Ser Ala Leu Lys 85 90 95

Gln Leu Asp Ile Gly Lys Val Val Phe Gly Cys Gly Asn Glu Arg Phe 100 105 110

Gly Gly Asn Gly Thr Val Leu Ser Val Asn His Asp Thr Cys Thr Leu 115 120 125

Val Pro Lys Asn Asn Ser Ala Ala Gly Tyr Glu Ser Ile Pro Gly Ile 130 135 140

Leu Arg Lys Glu Ala Ile Met Leu Leu Arg Tyr Phe Tyr Val Arg Gln
145 150 155 160

Asn Glu Arg Ala Pro Lys Pro Arg Ser Lys Ser Asp Arg Val Leu Asp 165 170 175

Lys Asn Thr Phe Pro Pro Met Glu Trp Ser Lys Tyr Leu Asn Glu Glu
180 185 190

Ala Phe Ile Glu Thr Phe Gly Asp Asp Tyr Arg Thr Cys Phe Ala Asn 195 200 205

Lys Val Asp Leu Ser Ser Asn Ser Val Asp Trp Asp Leu Ile Asp Ser 210 215 220

His Gln Asp Asn Ile Ile Gln Glu Leu Glu Glu Gln Cys Lys Met Phe 225 230 235 240

Lys Phe Asn Val His Lys Lys Ser Lys Val 245 250

<210> 147

<211> 3167

<212> DNA

<213> Saccharomyces cerevisiae

<400> 147

tcccctgatg gtcaaatact gtgcatggca tcccgtgcag tcaaggatgc tttgagacta 60 gttcatctgc cttcttgtag cgtgttcagc aactggccta ccagcgggac gcctttgggt 120

```
aaagttacca gtgtcgcatt ttcgccatct ggtgggctac tggccgtggg taacgaacaa 180
ggtaaagtga ggctctggaa attaaaccac tactaaattt ccatttatag acaaacttag 240
atattaaaag caatgtacaa atacatacac aaaatatcac tgtaaaaaaa ttgcgaagaa 300
acttgaaatt gaatatgatt ctgcccactt ttttcttgct gtcatttata gtcagaaatg 360
aaaaattgtc cgagaaatta aatataatat atggaaaaaa aggggacatt gagtttaaag 420
aatttgatta aaatgteete teaatateet etgtaagagt tatetaaate teaetttaet 480
tattcatctc gctgggaatt atgcaggcgg tagagagaag gccctcgtta ctctttgacg 540
aatatcaaaa ttctgttact aagcctaatg agacgaagaa taaagaagcc agggtcttgt 600
cagagaatga cggtgatgtc tccccatctg ttttgaaaca gaaggaaata tcagtcgatg 660
atatggatat gatttetttg cecaeggaat ttgacaggea aatggtttta ggttcaceta 720
tgtttttcga tcttgaagac gaagaaaaca aaattgatcc acttccttca gtttcccatc 780
attatggaaa tggagaaagt gacagctttg tctcatcgta cacgccctca aatctgaaaa 840
cgggtgaaga aactaaagat cttttcatta atccgtttga attggtttct caaatgagaa 900
aaagatacat tgctgcttcc aaacaagatg gcatttcaaa cataaaaaat gacactgaaa 960
agtggttttt atacccaaaa ccactgccaa agttttggag atttgaagac gataaacgat 1020
tccaagatcc ctctgactct gacttaaatg acgatggaga cagtactggg accggagccg 1080
ctacacegea cegecatgge tactattace caagttactt tacegateae tactactact 1140
acacaaaatc tggtttgaaa ggaaaaggaa atataaaagt accatacacc ggtgaatatt 1200
tcgatttaga ggattacaaa aaacaataca tttaccattt aagtaatcag gaaaatacgc 1260
aaaacccact ttcaccttat tctagtaagg aggagtcact agaggaagaa tttttaacag 1320
atgtgcctac gtttcaagaa tttagggatg attttgcata cataatagag ttaatccaat 1380
ctcataaatt caacgaggtt tcacgaaagc gattatctta tttattagat aaatttgaat 1440
tgtttcagta cctaaactct aagaaagaaa ttttagctaa taaaaatgtt ccctacagag 1500
atttttataa ttctcgtaag gtagatcgag acttgtcttt aagtggttgt atttctcaac 1560
gtcaattgag tgaatatata tgggagaaaa taaatttaga acctgaaagg atagtttatc 1620
aagacccgga aacgtcaagg aaactcagtt tgagagacat ttttcagttt ggttgttctt 1680
ctaatgacca acccattgcg attgggttga aattgattga tgatgaattc ttggattggt 1740
atagaaatat ttacctaata gattaccatc taactcctaa caaagtagca aagttggtcg 1800
gcaaagaaat gaggttttac ctattagcca aagtgtttct ggagtttgat aatttcattg 1860
aaggtgagta cctagcagaa attttcataa aatacgttat tcatatcctc gaaaaatcaa 1920
agtaccaatt ggcccaagta tcagttaatt ttcaattcta ttccagtggt gaagactggt 1980
acaagaaatt ttctcaatgg ttgctacgat ggaagctagt atcgtataat atccgctgga 2040
atatacaaat tgccaggatt tttcccaaac tattcaagga aaatgtcgtg tcaaatttcc 2100
aggagttttt ggatcttatc ttcaatcctt tattcactct ggaaaaggag cagttaccaa 2160
tagattcatc tgtaaatact gatatcattg gtctgcagtt ttttttatca aatgtgtgtt 2220
ctatggatct ggtcattaaa gagtcggatg aatattactg gaaagaattt actgatatga 2280
attgtaagcc aaaattttgg acagcacagg gtgacaatcc aactgttgcg cattacatgt 2340
attatattta taaaagttta gcgaaagtta attttctgcg gtcacaaaat cttcaaaata 2400
caatcaccct aagaaattat tgttctccac tatccagcag aacttcccaa tttggagtgg 2460
atttatattt tacagatcaa gttgaatcgt tagtgtgcaa cttactgctt tgtaatggtg 2520
gtctgctaca ggtagaaccg ctttgggata ctgcaacaat gattcaatat ttattttatc 2580
tettteaaat acceatttta getgegeeat tateatetgt tteattactg aattegeaaa 2640
aatcgacctt tttgaagaat aaaaacgtgc ttctagaaca tgattatttg aaagaccagg 2700
aaacagccaa aatcaatcct tctagagata tcactgtggg cgaacaaaga tcatatgaga 2760
caaatccttt catgaaaatg tttaagatgg gactaaaaat ttctttatca tcaaaatcga 2820
ttctttacaa tagttcatac acgctagaac ctctcattga agaatacagt gtagcagcaa 2880
gtatttactt gctgaaccca acagatttgt gcgagttgtc gagaacaagt gtgctatcta 2940
gtggctatga aggttggtac aaggctcatt ggattggcgt tggagttaaa aaggcgcctt 3000
```

actttgagga	gaacgtgggt	gggatagata	attggtacga	tacagcgaaa	gatacctcga	3060
taaagcacaa	cgttccgatg	attagaagaa	gatatagaaa	ggagacattg	gatcaagagt	3120
ggaacttcgt	tcgggatcac	tttggagtaa	ttaactccat	ttggtag		3167

<210> 148

<211> 888

<212> PRT

<213> Saccharomyces cerevisiae

<400> 148

Met Gln Ala Val Glu Arg Arg Pro Ser Leu Leu Phe Asp Glu Tyr Gln

1 5 10 15

Asn Ser Val Thr Lys Pro Asn Glu Thr Lys Asn Lys Glu Ala Arg Val 20 25 30

Leu Ser Glu Asn Asp Gly Asp Val Ser Pro Ser Val Leu Lys Gln Lys
35 40 45

Glu Ile Ser Val Asp Asp Met Asp Met Ile Ser Leu Pro Thr Glu Phe
50 55 60

Asp Arg Gln Met Val Leu Gly Ser Pro Met Phe Phe Asp Leu Glu Asp 65 70 75 80

Glu Glu Asn Lys Ile Asp Pro Leu Pro Ser Val Ser His His Tyr Gly 85 90 95

Asn Gly Glu Ser Asp Ser Phe Val Ser Ser Tyr Thr Pro Ser Asn Leu 100 105 110

Lys Thr Gly Glu Glu Thr Lys Asp Leu Phe Ile Asn Pro Phe Glu Leu 115 120 125

Val Ser Gln Met Arg Lys Arg Tyr Ile Ala Ala Ser Lys Gln Asp Gly 130 135 140

Ile Ser Asn Ile Lys Asn Asp Thr Glu Lys Trp Phe Leu Tyr Pro Lys 145 150 155 160

Pro Leu Pro Lys Phe Trp Arg Phe Glu Asp Asp Lys Arg Phe Gln Asp 165 170 175

Pro Ser Asp Ser Asp Leu Asn Asp Asp Gly Asp Ser Thr Gly Thr Gly 180 185 190

Ala Ala Thr Pro His Arg His Gly Tyr Tyr Tyr Pro Ser Tyr Phe Thr

<del>- 178 -</del>

200

- Asp His Tyr Tyr Tyr Tyr Thr Lys Ser Gly Leu Lys Gly Lys Gly Asn 210 215 220
- Ile Lys Val Pro Tyr Thr Gly Glu Tyr Phe Asp Leu Glu Asp Tyr Lys 225 230 235 240
- Lys Gln Tyr Ile Tyr His Leu Ser Asn Gln Glu Asn Thr Gln Asn Pro 245 250 255
- Leu Ser Pro Tyr Ser Ser Lys Glu Glu Ser Leu Glu Glu Glu Phe Leu 260 265 270
- Thr Asp Val Pro Thr Phe Gln Glu Phe Arg Asp Asp Phe Ala Tyr Ile 275 280 285
- Ile Glu Leu Ile Gln Ser His Lys Phe Asn Glu Val Ser Arg Lys Arg 290 295 300
- Leu Ser Tyr Leu Leu Asp Lys Phe Glu Leu Phe Gln Tyr Leu Asn Ser 305 310 315 320
- Lys Lys Glu Ile Leu Ala Asn Lys Asn Val Pro Tyr Arg Asp Phe Tyr 325 330 335
- Asn Ser Arg Lys Val Asp Arg Asp Leu Ser Leu Ser Gly Cys Ile Ser 340 345 350
- Gln Arg Gln Leu Ser Glu Tyr Ile Trp Glu Lys Ile Asn Leu Glu Pro 355 360 365
- Glu Arg Ile Val Tyr Gln Asp Pro Glu Thr Ser Arg Lys Leu Ser Leu 370 375 380
- Arg Asp Ile Phe Gln Phe Gly Cys Ser Ser Asn Asp Gln Pro Ile Ala 385 390 395 400
- Ile Gly Leu Lys Leu Ile Asp Asp Glu Phe Leu Asp Trp Tyr Arg Asn 405 410 415
- Ile Tyr Leu Ile Asp Tyr His Leu Thr Pro Asn Lys Val Ala Lys Leu 420 425 430
- Val Gly Lys Glu Met Arg Phe Tyr Leu Leu Ala Lys Val Phe Leu Glu
  435 440 445
- Phe Asp Asn Phe Ile Glu Gly Glu Tyr Leu Ala Glu Ile Phe Ile Lys

UL UI 1333

460

450 455

Tyr Val Ile His Ile Leu Glu Lys Ser Lys Tyr Gln Leu Ala Gln Val 465 470 475 480

Ser Val Asn Phe Gln Phe Tyr Ser Ser Gly Glu Asp Trp Tyr Lys Lys 485 490 495

Phe Ser Gln Trp Leu Leu Arg Trp Lys Leu Val Ser Tyr Asn Ile Arg 500 505 510

Trp Asn Ile Gln Ile Ala Arg Ile Phe Pro Lys Leu Phe Lys Glu Asn 515 520 525

Val Val Ser Asn Phe Gln Glu Phe Leu Asp Leu Ile Phe Asn Pro Leu 530 535 540

Phe Thr Leu Glu Lys Glu Gln Leu Pro Ile Asp Ser Ser Val Asn Thr 545 550 555 560

Asp Ile Ile Gly Leu Gln Phe Phe Leu Ser Asn Val Cys Ser Met Asp 565 570 575

Leu Val Ile Lys Glu Ser Asp Glu Tyr Tyr Trp Lys Glu Phe Thr Asp 580 585 590

Met Asn Cys Lys Pro Lys Phe Trp Thr Ala Gln Gly Asp Asn Pro Thr 595 600 605

Val Ala His Tyr Met Tyr Tyr Ile Tyr Lys Ser Leu Ala Lys Val Asn 610 615 620

Phe Leu Arg Ser Gln Asn Leu Gln Asn Thr Ile Thr Leu Arg Asn Tyr 625 630 635 640

Cys Ser Pro Leu Ser Ser Arg Thr Ser Gln Phe Gly Val Asp Leu Tyr
645 650 655

Phe Thr Asp Gln Val Glu Ser Leu Val Cys Asn Leu Leu Cys Asn 660 665 670

Gly Gly Leu Leu Gln Val Glu Pro Leu Trp Asp Thr Ala Thr Met Ile 675 680 685

Gln Tyr Leu Phe Tyr Leu Phe Gln Ile Pro Ile Leu Ala Ala Pro Leu 690 695 700

Ser Ser Val Ser Leu Leu Asn Ser Gln Lys Ser Thr Phe Leu Lys Asn

705 710 715 720

Lys Asn Val Leu Leu Glu His Asp Tyr Leu Lys Asp Gln Glu Thr Ala 725 730 735

Lys Ile Asn Pro Ser Arg Asp Ile Thr Val Gly Glu Gln Arg Ser Tyr 740 745 750

Glu Thr Asn Pro Phe Met Lys Met Phe Lys Met Gly Leu Lys Ile Ser 755 760 765

Leu Ser Ser Lys Ser Ile Leu Tyr Asn Ser Ser Tyr Thr Leu Glu Pro 770 775 780

Leu Ile Glu Glu Tyr Ser Val Ala Ala Ser Ile Tyr Leu Leu Asn Pro 785 790 795 800

Thr Asp Leu Cys Glu Leu Ser Arg Thr Ser Val Leu Ser Ser Gly Tyr 805 810 815

Glu Gly Trp Tyr Lys Ala His Trp Ile Gly Val Gly Val Lys Lys Ala 820 825 830

Pro Tyr Phe Glu Glu Asn Val Gly Gly Ile Asp Asn Trp Tyr Asp Thr 835 840 845

Ala Lys Asp Thr Ser Ile Lys His Asn Val Pro Met Ile Arg Arg Arg 850 855 860

Tyr Arg Lys Glu Thr Leu Asp Gln Glu Trp Asn Phe Val Arg Asp His 865 870 875 880

Phe Gly Val Ile Asn Ser Ile Trp 885

<210> 149

<211> 3146

<212> DNA

<213> Saccharomyces cerevisiae

<400> 149

tttttcttcg cacatggctg gttctgctgg aggcggctgg gtgtttatag ttaccgcaat 60 gcaggtcgct tcgtttactt ttctctgcgt gctcgagaac tgataaatac tggtttagga 120 aaaccgtcca cttatggatt taacaaggag gtgaaaggta ggtaccatgg caatatagct 180 agcaacgcta cgcactagct gtgtgtgctt gctctcttgt gttcgtcttt cgtatattgc 240 gggttcccgc ggttaattca acaacaata tggtatctag gcagggaccc agcaacggaa 300

ggatgtaatt	gctatatagg	gccgatgaaa	cagttctcat	gtttcaactc	ggtcgaagaa	360
ttctctaaaa	catattctat	acttcaaagt	ttttagcttc	ttgcttttcg	gtagttgcca	420
aaccaatatc	ggcataaaat	tataattgct	gcacccctgc	aactccaaac	aacttaaaac	480
aacttaaaac	gaaaacgctt	atgctggagt	ttccaatatc	agttctgcta	ggatgcctag	540
					aacgagcata	
					actctggcca	
					cattccgatg	
					gacgcctggt	
					acgggtcact	
					tattgtggta	
					ctgggtgagt	
					tcgtcctctt	
					cccgctgtag	
					aaatcgacga	
					attactagta	
					gtcactactt	
					actaccacca	
					agcactgctg	
					gcgagtaccg	
					tctgctcctt	
					aattcagtta	
					tctgtaagcg	
					aaggaggcca	
					gcccaatatg	
					tcagctgtgt	
cgcaatctgt	tctgaattcc	gttatagccg	tcaacaccga	cgtatctgta	acctcagtta	1860
gtagcacagc	ccataccaca	aaggacaccg	ccaccacttc	agtaaccgcc	tcagaaagta	1920
tcacttcgga	aactgctcag	gcttcaagtt	caacagagaa	gaatattagt	aacagtgccg	1980
ccacatcgag	tagcatttac	tccaacagtg	cttctgtgtc	aggacacggt	gtaacatacg	2040
ctgccgaata	cgccattaca	tccgagcaat	cctctgcgct	tgccacatct	gtgcctgcta	2100
					acaaccatca	
					acaagggcag	
					agtccaaccg	
					ttagatagct	
					tctacctgca	
					accaccactg	
					aatttcccta	
					acatcttgta	
					atgaaaacca	
					gaaaatgatg	
					aaccccggag	
					cccacttctg	
					cagactgttg	
					actggtagca	
					acttcctcga	
					tacgaaggtg	
			atagtctagc	tggaacgatt	gccgcttttg	3120
tcgtagctgt	tctgttcgcc	ttctag				3146

<210> 150

<211> 881

<212> PRT

<213> Saccharomyces cerevisiae

<400> 150

Met Leu Glu Phe Pro Ile Ser Val Leu Leu Gly Cys Leu Val Ala Val 1 5 10 15

Lys Ala Gln Thr Thr Phe Pro Asn Phe Glu Ser Asp Val Leu Asn Glu 20 25 30

His Asn Lys Phe Arg Ala Leu His Val Asp Thr Ala Pro Leu Thr Trp 35 40 45

Ser Asp Thr Leu Ala Thr Tyr Ala Gln Asn Tyr Ala Asp Gln Tyr Asp 50 55 60

Cys Ser Gly Val Leu Thr His Ser Asp Gly Pro Tyr Gly Glu Asn Leu 65 70 75 80

Ala Leu Gly Tyr Thr Asp Thr Gly Ala Val Asp Ala Trp Tyr Gly Glu 85 90 95

Ile Ser Lys Tyr Asn Tyr Ser Asn Pro Gly Phe Ser Glu Ser Thr Gly
100 105 110

His Phe Thr Gln Val Val Trp Lys Ser Thr Ala Glu Ile Gly Cys Gly 115 120 125

Tyr Lys Tyr Cys Gly Thr Thr Trp Asn Asn Tyr Ile Val Cys Ser Tyr 130 135 140

Asn Pro Pro Gly Asn Tyr Leu Gly Glu Phe Ala Glu Glu Val Glu Pro 145 150 155 160

Leu Ile Ser Thr Val Ser Ser Ser Ser Ser Ser Ser Ser Ser Thr Ser 165 170 175

Thr Thr Ser Asp Thr Val Ser Thr Ile Ser Ser Ser Ile Met Pro Ala 180 185 190

Val Ala Gln Gly Tyr Thr Thr Thr Val Ser Ser Ala Ala Ser Ser Ser 195 200 205

Ser Leu Lys Ser Thr Thr Ile Asn Pro Ala Lys Thr Ala Thr Leu Thr

215

220

Ala Ser Ser Ser Thr Val Ile Thr Ser Ser Thr Glu Ser Val Gly Ser 225 230 235 240

Ser Thr Val Ser Ser Ala Ser Ser Ser Ser Val Thr Thr Ser Tyr Ala 245 250 255

Thr Ser Ser Ser Thr Val Val Ser Ser Asp Ala Thr Ser Ser Thr Thr 260 265 270

Thr Thr Ser Ser Val Ala Thr Ser Ser Ser Thr Thr Ser Ser Asp Pro 275 280 285

Thr Ser Ser Thr Ala Ala Ala Ser Ser Ser Asp Pro Ala Ser Ser Ser 290 295 300

Ala Ala Ser Ser Ser Ala Ser Thr Glu Asn Ala Ala Ser Ser Ser 305 310 315 320

Ser Ala Ile Ser Ser Ser Ser Met Val Ser Ala Pro Leu Ser Ser 325 330 335

Thr Leu Thr Thr Ser Thr Ala Ser Ser Arg Ser Val Thr Ser Asn Ser 340 345 350

Val Asn Ser Val Lys Phe Ala Asn Thr Thr Val Phe Ser Ala Gln Thr 355 360 365

Thr Ser Ser Val Ser Ala Ser Leu Ser Ser Ser Val Ala Ala Asp Asp 370 375 380

Ile Gln Gly Ser Thr Ser Lys Glu Ala Thr Ser Ser Val Ser Glu His 385 390 395 400

Thr Ser Ile Val Thr Ser Ala Thr Asn Ala Ala Gln Tyr Ala Thr Arg
405 410 415

Leu Gly Ser Ser Ser Ser Ser Ser Gly Ala Val Ser Ser Ser Ala
420 425 430

Val Ser Gln Ser Val Leu Asn Ser Val Ile Ala Val Asn Thr Asp Val 435 440 445

Ser Val Thr Ser Val Ser Ser Thr Ala His Thr Thr Lys Asp Thr Ala 450 455 460

Thr Thr Ser Val Thr Ala Ser Glu Ser Ile Thr Ser Glu Thr Ala Gln

465					470					475		•			480
Ala	Ser	Ser	Ser	Thr 485	Glu	Lys	Asn	Ile	Ser 490	Asn	Ser	Ala	Ala	Thr 495	Ser
Ser	Ser	Ile	Tyr 500	Ser	Asn	Ser	Ala	Ser 505	Val	Ser	Gly	His	Gly 510	Val	Thr
Tyr	Ala	Ala 515	Glu	Туг	Ala	Ile	Thr 520	Ser	Glu	Gln	Ser	Ser 525	Ala	Leu	Ala
Thr	Ser 530	Val	Pro	Ala	Thr	Asn 535	Cys	Ser	Ser	Ile	Val 540	Lys	Thr	Thr	Thr
Leu 545	Glu	Asn	Ser	Ser	Thr 550	Thr	Thr	Ile	Thr	Ala 555	Ile	Thr	Lys	Ser	Thr 560
Thr	Thr	Leu	Ala	Thr 565		Ala	Asn	Asn	Ser 570	Thr	Arg	Ala	Ala	Thr 575	Ala
Val	Thr	Ile	Asp 580		Thr	Leu	. Asp	Pro 585		Asp	Asn	Ser	Ala 590	Ser	Pro
Thr	Asp	Asn 595		Lys	His	Thr	Ser 600		Туг	: Gly	Ser	Ser 605	Ser	Thr	Gly
Ala	Ser		ı Asp	Ser	Leu	Arg 615		Thr	Thr	Ser	1le 620	Ser	Va]	Ser	Ser
Asn 625		Thr	Glr	. Lev	Val		Thr	Cys	3 Thi	635	Glu 5	Sei	Ası	у Туі	640
Asp	Ser	Pro	Sei	Phe 645		ı Ile	e Sei	Thi	650	a Thi	r Thi	Thi	r Gli	u Se:	r Asn 5
Lev	ı Ile	e Thi	r Ası 660		r Ile	e Thi	r Ala	66		s Se	r Thi	c As	p Se 67	r As: O	n Phe
Pro	Th:	r Se:		a Ala	a Sei	r Se:	r Thi		p Gl	u Th	r Ala	a Ph 68	e Th 5	r Ar	g Thr
Ile	e Se:		r Se	r Cy	s Se:	r Th		u As	n Gl	y Al	a Se:	r Th O	r G1	n Th	r Ser

710

Glu Leu Thr Thr Ser Pro Met Lys Thr Asn Thr Val Val Pro Ala Ser

Ser Phe Pro Ser Thr Thr Thr Cys Leu Glu Asn Asp Asp Thr Ala

715

725 730 735

Phe Ser Ser Ile Tyr Thr Glu Val Asn Ala Ala Thr Ile Ile Asn Pro
740 745 750

Gly Glu Thr Ser Ser Leu Ala Ser Asp Phe Ala Thr Ser Glu Lys Pro 755 760 765

Asn Glu Pro Thr Ser Val Lys Ser Thr Ser Asn Glu Gly Thr Ser Ser 770 775 780

Thr Thr Thr Tyr Gln Gln Thr Val Ala Thr Leu Tyr Ala Lys Pro 785 790 795 800

Ser Ser Thr Ser Leu Gly Ala Arg Thr Thr Thr Gly Ser Asn Gly Arg 805 810 815

Ser Thr Thr Ser Gln Gln Asp Gly Ser Ala Met His Gln Pro Thr Ser 820 825 830

Ser Ile Tyr Thr Gln Leu Lys Glu Gly Thr Ser Thr Thr Ala Lys Leu 835 840 845

Ser Ala Tyr Glu Gly Ala Ala Thr Pro Leu Ser Ile Phe Gln Cys Asn 850 855 860

Ser Leu Ala Gly Thr Ile Ala Ala Phe Val Val Ala Val Leu Phe Ala 865 870 875 880

Phe

<210> 151

<211> 830

<212> DNA

<213> Saccharomyces cerevisiae

<400> 151

ttgcattatc aactataatt ccattccta gaggagttt tatagtcctg ccatctaatt 60 gtagagcaat ctttccttc tcaacatccc tatttagtga caccttttcc caaaacttct 120 gtgaagtttt gctcaatcta tttgtttccg taggagtgt attctaatc gtgttgtctg 180 tccccagtgg ctgggcattt aatgagtaga atcggggcag tttcaatctt atggaattca 240 caatgaagca tcccttcctt aatgatggca gcattccgta ttctcttttg tagggtttcg 300 tttgccttca agtgtttgt tcttatttag cctttcctt taccttaatt tttttctttt 360 ctctgaagaa aatgaatgag tttaaagata tagcaattaa aaagtaacag tgaagaaatt 420 tctcagatga gcagatggga attaaagaac tatctacaga gctctttact aaattgaatc 480



```
aataatacat acttacaaac atgtcacaga tagcacaaga aatgacagtg agcttaagaa 540
acgccaggac acaattggat atggtcaatc agcagctagc atatttggac agacaagaaa 600
agettgetga attgacaaag aaagaactag agtettatee aaeggacaaa gtatggagat 660
cttgcggtaa atcgtttatc ttacaggata aatccaaata cgttaatgat ttatcacatg 720
ccgaaactgt tcttctggat caaagaaaaa cattaaagat aaagaagaac tatttagaaa 780
ctactgttga aaaaacaata gacaatctaa aggcattgat gaagaattaa
                                                                   830
<210> 152
<211> 109
<212> PRT
<213> Saccharomyces cerevisiae
<400> 152
Met Ser Gln Ile Ala Gln Glu Met Thr Val Ser Leu Arg Asn Ala Arg
Thr Gln Leu Asp Met Val Asn Gln Gln Leu Ala Tyr Leu Asp Arg Gln
             20
                                 25
Glu Lys Leu Ala Glu Leu Thr Lys Lys Glu Leu Glu Ser Tyr Pro Thr
         35
                             40
                                                  45
Asp Lys Val Trp Arg Ser Cys Gly Lys Ser Phe Ile Leu Gln Asp Lys
  · 50
                         55
Ser Lys Tyr Val Asn Asp Leu Ser His Ala Glu Thr Val Leu Leu Asp
                     70
                                          75
Gln Arg Lys Thr Leu Lys Ile Lys Lys Asn Tyr Leu Glu Thr Thr Val
Glu Lys Thr Ile Asp Asn Leu Lys Ala Leu Met Lys Asn
<210> 153
<211> 1478
<212> DNA
<213> Saccharomyces cerevisiae
<400> 153
aatagttctt ctttatcttt aatgtttttc tttgatccag aagaacagtt tcggcatgtg 60
```

ataaatcatt aacgtatttg gatttatcct gtaagataaa cgatttaccg caagatctcc 120 atactttgtc cgttggataa gactctagtt ctttctttgt caattcagca agcttttctt 180 gtctgtccaa atatgctagc tgctgattga ccatatccaa ttgtgtcctg gcgtttctta 240 agctcactgt catttcttgt gctatctgtg acatgtttgt aagtatgtat tattgattca 300

```
atttagtaaa gagctctgta gatagttctt taattcccat ctgctcatct gagaaatttc 360
ttcactgtta ctttttaatt gctatatctt taaactcatt cattttcttc agagaaaaga 420
aaaaaattaa ggtaaaggaa aaggctaaat aagaacaaaa cacttgaagg caaacgaaac 480
cctacaaaag agaatacgga atgctgccat cattaaggaa gggatgcttc attgtgaatt 540
ccataagatt gaaactgccc cgattctact cattaaatgc ccagccactg gggacagaca 600
acacgattga gaataacact cctacggaaa caaatagatt gagcaaaact tcacagaagt 660
tttgggaaaa ggtgtcacta aatagggatg ttgagaaagg aaagattgct ctacaattag 720
atggcaggac tataaaaact cctctaggaa atggaattat agttgataat gcaaagtctc 780
tettageata cetattaaaa etggagtggt egteectate eagtetttee ateaaaacte 840
actetttgcc actaacttca ttagtggcaa gatgcataga tttacaaatg acaaatgagc 900
ctggctgtga ccctcaatta gttgcaaaga ttggaggcaa cagtgatgtt ataaaaaatc 960
agttgttaag atatttagat accgatactt tattggtctt ttcccctatg aatgagtttg 1020
aaggaagatt acgcaatgcg caaaatgagt tatatatacc catcatcaaa ggaatggaag 1080
agtttttacg caacttttca tccgagtcta atattcgact acaaatttta gatgccgaca 1140
tccatgggtt acgaggcaat cagcagtcgg atatcgttaa gaatgcagca aaaaaatata 1200
tgagcagctt atcaccatgg gatcttgcaa ttcttgaaaa aactgtatta accacaaagt 1260
ccttcatttg cggcgtgcta ttattagaaa ataaaaaaga tactgcgaac ttaattcccg 1320
ccttgaaaac tgatatggat aatattgtac gtgccgccac cttagaaaca atcttccaag 1380
ttgaaaagtg gggagaggtt gaagatactc atgacgttga caaaagagac atcagaagaa 1440
                                                                  1478
aaattcatac tgctgcgatt gctgctttta agcaataa
```

```
<210> 154
```

<211> 325

<212> PRT

<213> Saccharomyces cerevisiae

<400> 154

Met Leu Pro Ser Leu Arg Lys Gly Cys Phe Ile Val Asn Ser Ile Arg 1 5 10 15

Leu Lys Leu Pro Arg Phe Tyr Ser Leu Asn Ala Gln Pro Leu Gly Thr
20 25 30

Asp Asn Thr Ile Glu Asn Asn Thr Pro Thr Glu Thr Asn Arg Leu Ser 35 40 45

Lys Thr Ser Gln Lys Phe Trp Glu Lys Val Ser Leu Asn Arg Asp Val 50 55 60

Glu Lys Gly Lys Ile Ala Leu Gln Leu Asp Gly Arg Thr Ile Lys Thr
65 70 75 80

Pro Leu Gly Asn Gly Ile Ile Val Asp Asn Ala Lys Ser Leu Leu Ala 85 90 95

Tyr Leu Leu Lys Leu Glu Trp Ser Ser Leu Ser Ser Leu Ser Ile Lys 100 105 110

Thr His Ser Leu Pro Leu Thr Ser Leu Val Ala Arg Cys Ile Asp Leu 115 120 125

Gln Met Thr Asn Glu Pro Gly Cys Asp Pro Gln Leu Val Ala Lys Ile 130 135 140

Gly Gly Asn Ser Asp Val Ile Lys Asn Gln Leu Leu Arg Tyr Leu Asp 145 150 155 160

Thr Asp Thr Leu Leu Val Phe Ser Pro Met Asn Glu Phe Glu Gly Arg 165 170 175

Leu Arg Asn Ala Gln Asn Glu Leu Tyr Ile Pro Ile Ile Lys Gly Met
180 185 190

Glu Glu Phe Leu Arg Asn Phe Ser Ser Glu Ser Asn Ile Arg Leu Gln 195 200 205

Ile Leu Asp Ala Asp Ile His Gly Leu Arg Gly Asn Gln Gln Ser Asp 210 215 220

Ile Val Lys Asn Ala Ala Lys Lys Tyr Met Ser Ser Leu Ser Pro Trp 225 230 235 240

Asp Leu Ala Ile Leu Glu Lys Thr Val Leu Thr Thr Lys Ser Phe Ile 245 250 255

Cys Gly Val Leu Leu Leu Glu Asn Lys Lys Asp Thr Ala Asn Leu Ile 260 265 270

Pro Ala Leu Lys Thr Asp Met Asp Asn Ile Val Arg Ala Ala Thr Leu 275 280 285

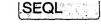
Glu Thr Ile Phe Gln Val Glu Lys Trp Gly Glu Val Glu Asp Thr His 290 295 300

Asp Val Asp Lys Arg Asp Ile Arg Arg Lys Ile His Thr Ala Ala Ile 305 310 315 320

Ala Ala Phe Lys Gln 325

<210> 155
<211> 2336
<212> DNA





## <213> Saccharomyces cerevisiae

```
<400> 155
gcaagtcagt acgatgcatc cttggatcaa gaaggtgagt ctggaaacgg tgcttctaat 60
ggcgatgttt accattataa tgaaggggac ttggctgcct ccttcaaggg atgtagatca 120
agaggtacgt gtgctagtga aataggtcac atgtaccaga aaatcaagaa atcttagatg 180
tgtgtaatct tgcaatttag tttcaaacaa gatccttgta tttatataaa caaagatata 240
atttttaaaa aaaaattaga aaaagcaaat ataattcagg tcccacttgg aataatggca 300
ctgtattgat gcattttcct tatgcttagt gacgcgtttt cgcgcgtcag tttcaagttt 360
ttcttggctt tttttttttc attttcgtaa agggtcttaa aaggattaaa aaatgcagta 420
ttgaaataaa gaacaattac gaacggtgaa gctgctattt tggttattat acccttccag 480
gacagtacgc gcaaactatt atggagatat tcaaggaaga agaagaagaa gctttttcgg 540
cgatagaagg tataatatat gcctgtgagg tgtatgaccc tgtaccccgt catttacata 600
aaagcaaaac aaagatcatc aatgctgcta aattaattat agaaacgcat ctttcatatt 660
atacaatact caataacatt tcagatatac aagcctatct ttctacttgg cttagggatc 720
ttggaacgac aggtccatac caaacaattc tttcagaaag tatttctctc atgtttgacc 780
gcactgtatc tatcttcagg aaatgtacga tagagggagg ttttccacat ttgatcgcac 840
gtctttatct cagattgaaa agctaccaga agctcttaaa cgatgcagga ttaaagaatt 900
ttttttcaag ctacgattat gctttcgggg ttgcatacaa ccttgtaaat tgctctgaat 960
acaggtatga cgaagttcat tacatatcga acggcactta ctcattagtt gcatcgatga 1020
agatagatcc tgctgaagtc attaaaaggg aacattttag gcttacaatt ccgaaattta 1080
acatatctaa tatattaatt gaaatttttc atttgctcga tggattagca ttttttaagg 1140
tgaaccctga tagtttatct atatctacag cttcagcaga aacaatcttt cgcagtatct 1200
ccgaaggtaa tcatcaggtc ctagaattgg ggagaagttt aatgtttcca ttgttgagga 1260
ctggagattt tgaaatctgt cgtattgacg acgcgggagc tgtcataaca tttacagaag 1320
cgaaggatgt aaaactagaa ataatcagtc tggatgaagt ttcctgggta atgcagtgga 1380
aatcttgtct tcaaaattat gagagaaggg cagcaaatga cagttcattt atcaaaacac 1440
acctacaatt taagaaggcc aacaatttca atgaagataa taatgggcta ggactaattg 1500
tagacagaaa tattecaaca gatgatttta egetagette tacaaacegt caaagteece 1560
cgccttcaaa tactggttgt tcattacaca ggtctaaacc cttgcatatc cctttatcat 1620
ctgttattcg tgaagacttt tatgatagct ctctaaatga gcgtatatct aaagacggag 1680
atagcagttg tgaatccttc agtggcgccg aaagtatctt atcagactac gattttcatg 1740
ataatgaatt ttttaacaac cagtcacctc attatttttc agaacacata gacaataact 1800
cgagagaggt ggtaataaca gatgaaaata cgataatatc tttggaaaat acccaagtaa 1860
gtcggtggtc aaattactca tggcaaaaaa tttcaccgca tcaattacag gtctctatta 1920
tccaactgcg catgggaaac ttcattgtgg cttatgattc tgattataac cttcatcagt 1980
tcaaaattcg tttgtgtgac gatataaaat gtatacaatc cacagagcaa gacatacaaa 2040
tacgtgtccc gctcggcgca ataatgtgca gcgtcactgg tatcttgaat attaggacga 2100
aggacgctga caagttgctt cgggtattaa gcttttatac cactgaccac acggaagctg 2160
tatcgcactc aaacaatcaa gatgctactg caagtccact ttcgtcagtt tcatcagcaa 2220
tggateteaa geatteatta cagaaatgtt eetetaeaat aatgeeecaa gagttgaege 2280
aggacgtcat cggttcaaaa tcagacctaa tcagtaatat tcgtcaaaaa atataa
                                                                  2336
```

<210> 156

<211> 611

<212> PRT

<213> Saccharomyces cerevisiae

-1	Λ	^	1	5	f

Met Glu Ile Phe Lys Glu Glu Glu Glu Glu Ala Phe Ser Ala Ile Glu 5

Gly Ile Il Tyr Ala Cys Glu Val Tyr Asp Pro Val Pro Arg His Leu 25

His Lys Ser Lys Thr Lys Ile Ile Asn Ala Ala Lys Leu Ile Ile Glu 40 35

Thr His Leu Ser Tyr Tyr Thr Ile Leu Asn Asn Ile Ser Asp Ile Gln 55 50

Ala Tyr Leu Ser Thr Trp Leu Arg Asp Leu Gly Thr Thr Gly Pro Tyr 75 70

Gln Thr Ile Leu Ser Glu Ser Ile Ser Leu Met Phe Asp Arg Thr Val 85

Ser Ile Phe Arg Lys Cys Thr Ile Glu Gly Gly Phe Pro His Leu Ile 110 105 100

Ala Arg Leu Tyr Leu Arg Leu Lys Ser Tyr Gln Lys Leu Leu Asn Asp 120 115

Ala Gly Leu Lys Asn Phe Phe Ser Ser Tyr Asp Tyr Ala Phe Gly Val 135 130

Ala Tyr Asn Leu Val Asn Cys Ser Glu Tyr Arg Tyr Asp Glu Val His 155 150 145

Tyr Ile Ser Asn Gly Thr Tyr Ser Leu Val Ala Ser Met Lys Ile Asp 170 165

Pro Ala Glu Val Ile Lys Arg Glu His Phe Arg Leu Thr Ile Pro Lys 185

Phe Asn Ile Ser Asn Ile Leu Ile Glu Ile Phe His Leu Leu Asp Gly 200 195

Leu Ala Phe Phe Lys Val Asn Pro Asp Ser Leu Ser Ile Ser Thr Ala 220 215 210

Ser Ala Glu Thr Ile Phe Arg Ser Ile Ser Glu Gly Asn His Gln Val 235 230 225

Leu Glu Leu Gly Arg Ser Leu Met Phe Pro Leu Leu Arg Thr Gly Asp



255

Phe Glu Ile Cys Arg Ile Asp Asp Ala Gly Ala Val Ile Thr Phe Thr 260 265 270

Glu Ala Lys Asp Val Lys Leu Glu Ile Ile Ser Leu Asp Glu Val Ser 275 280 285

Trp Val Met Gln Trp Lys Ser Cys Leu Gln Asn Tyr Glu Arg Arg Ala 290 295 300

Ala Asn Asp Ser Ser Phe Ile Lys Thr His Leu Gln Phe Lys Lys Ala 305 310 315 320

Asn Asn Phe Asn Glu Asp Asn Asn Gly Leu Gly Leu Ile Val Asp Arg
325 330 335

Asn Ile Pro Thr Asp Asp Phe Thr Leu Ala Ser Thr Asn Arg Gln Ser 340 345 350

Pro Pro Pro Ser Asn Thr Gly Cys Ser Leu His Arg Ser Lys Pro Leu 355 360 365

His Ile Pro Leu Ser Ser Val Ile Arg Glu Asp Phe Tyr Asp Ser Ser 370 380

Leu Asn Glu Arg Ile Ser Lys Asp Gly Asp Ser Ser Cys Glu Ser Phe 385 390 395 400

Ser Gly Ala Glu Ser Ile Leu Ser Asp Tyr Asp Phe His Asp Asn Glu
405 410 415

Phe Phe Asn Asn Gln Ser Pro His Tyr Phe Ser Glu His Ile Asp Asn 420 425 430

Asn Ser Arg Glu Val Val Ile Thr Asp Glu Asn Thr Ile Ile Ser Leu 435 440 445

Glu Asn Thr Gln Val Ser Arg Trp Ser Asn Tyr Ser Trp Gln Lys Ile 450 455 460

Ser Pro His Gln Leu Gln Val Ser Ile Ile Gln Leu Arg Met Gly Asn 465 470 475 480

Phe Ile Val Ala Tyr Asp Ser Asp Tyr Asn Leu His Gln Phe Lys Ile
485 490 495

Arg Leu Cys Asp Asp Ile Lys Cys Ile Gln Ser Thr Glu Gln Asp Ile

EP99870141.1

SÉQL

500 505 510

Gln Ile Arg Val Pro Leu Gly Ala Ile Met Cys Ser Val Thr Gly Ile 515 520 525

Leu Asn Ile Arg Thr Lys Asp Ala Asp Lys Leu Leu Arg Val Leu Ser 530 535 540

Phe Tyr Thr Thr Asp His Thr Glu Ala Val Ser His Ser Asn Asn Gln 545 550 555 560

Asp Ala Thr Ala Ser Pro Leu Ser Ser Val Ser Ser Ala Met Asp Leu 565 570 575

Lys His Ser Leu Gln Lys Cys Ser Ser Thr Ile Met Pro Gln Glu Leu 580 585 590

Thr Gln Asp Val Ile Gly Ser Lys Ser Asp Leu Ile Ser Asn Ile Arg 595 600 605

Gln Lys Ile 610

<210> 157

<211> 2960

<212> DNA

<213> Saccharomyces cerevisiae

<400> 157

tcttgcaacc atcgtcctct aaggaaagaa atgtggcacc tttaaatagg ttccacctca 60 cagatgccaa catgtgggag atagggggct attcgcaatt tattaacgtc tctagtactg 120 gtaagccttt ctaacattct tgtggtcacg tgatgtgtat gtttcttttt tttattcctc 180 cgggtgataa cgacgcgaaa aatattttt catttttctt ttaaccatca caatttgcga 240 cgcgacgcga aaaaaatgca gagaagtaaa agaatgctgg ataagaaatc aacaacacag 300 ttgaacattg gcgtgcccct gtgtattaag taaaggttgt ccaagaggct cttttttgta 360 tgtatgcgtg tgttaactat cctgcacatc atcttgcgca gttagtccaa taaaaaagga 420 ttactactga acaggtetta etatttttga ttgcgtagtg etggggaaaa gtaaacacac 480 acaggcgcac acgagaacag atgagttctt tggacgagga tgaagaggac ttcgaaatgc 540 tggacacgga gaacctccag tttatgggga agaagatgtt tggcaaacag gccggcgaag 600 acgagagtga tgattttgct atagggggta gcaccccgac caataaactg aaattttatc 660 catattcgaa caacaaattg acaagaagta cggggacctt gaacctgtca ttaagtaata 720 cagctttgtc agaggctaac tccaaatttc ttgggaaaat tgaagaagag gaagaagagg 780 aggaagaagg caaggatgag gaaagcgtgg attctcgtat taaaaggtgg tctccgttcc 840 atgaaaatga aagtgttact actcctatta caaaaagatc tgcggaaaaa acgaacagtc 900 ctatttctct caaacaatgg aaccagcgat ggtttccgaa aaatgatgct cgcactgaaa 960 atacatecte atectettea tatagegteg etaaacetaa ecaateagee titaegtett 1020

```
cgggcctcgt atctaaaatg tctatggaca cttcgttata ccctgcgaaa ttgaggatac 1080
cagaaacacc agtgaaaaaa tcacccttag tggagggaag agaccataag catgtccacc 1140
tttcgagttc gaaaaatgca tcgtcttctc taagtgtttc ccctttaaat tttgttgaag 1200
acaataattt acaagaagac cttttatttt cagattctcc gtcttcgaaa gctttacctt 1260
ccatccatgt accaaccata gactcatccc cactgagcga ggcaaaatat catgcacatg 1320
atogtoacaa taaccagaca aacatootgt otoccactaa tagottggtt accaacagot 1380
ctccacaaac attgcattct aacaagttca aaaaaatcaa aagagcaagg aattcggtta 1440
ttttgaaaaa tagagagcta acaaacagtt tacaacaatt caaagatgat ttatacggca 1500
cggacgagaa tttcccacct ccaatcataa tatcaagtca tcattcaact agaaagaacc 1560
ctcaacctta tcaatttcgt ggacgctatg acaatgacac tgacgaagag atctccactc 1620
caacaagacg aaaatctatt attggggcaa catctcaaac acatagagaa agcagaccat 1680
tgtcactctc ctctgccatc gtgacaaaca caacaagtgc agagacgcat tccatatctt 1740
ccaccgattc ttcgccgtta aattccaaaa ggcgtctaat ctcttcaaat aagttatcag 1800
caaatccaga ttcccatctt ttcgaaaaat ttacgaatgt gcattccatt ggtaaaggcc 1860
agttttccac ggtctaccag gttacgtttg cccaaacaaa caaaaagtat gcaatcaaag 1920
ccattaaacc aaacaaatat aattccttga aacgcatatt actggaaatt aaaatactaa 1980
acgaggtaac aaaccaaatt acaatggatc aagaagggaa ggaatacatc atcgattaca 2040
tragttrictg gaagtttraa aattratact atattatgar agaattgtgr gaaaatggta 2100
atttggatgg atttttacaa gagcaagtta tcgcaaagaa aaaaaggttg gaagattgga 2160
gaatttggaa aatcatcgtg gaattaagcc tggctttacg attcatccat gattcttgtc 2220
acattgtgca tctggacttg aaacccgcaa acgtcatgat cacatttgaa ggtaacctaa 2280
aactaggtga ctttggaatg gctactcatt taccgttgga ggataaaagt tttgaaaatg 2340
aaggtgacag agaatatatt gcaccagaaa tcatttctga ttgtacgtac gattacaagg 2400
cagatatttt ttccctgggt ctgatgattg ttgaaattgc agcgaacgtt gtgttacctg 2460
acaatggcaa cgcatggcat aagttgagat cgggtgattt atcggatgca ggaagattaa 2520
gttccacaga tattcattct gaatcattat tttcagacat tacgaaagta gatacaaatg 2580
atttatttga ttttgaaaga gacaatatca gtggtaatag taacaacgct`ggcacctcca 2640
ctgttcataa caatagtaat atcaacaacc ctaatatgaa taatggcaac gataataata 2700
atgtcaatac tgccgctacc aagaatcgtc ttattttgca taaaagttct aaaattcccg 2760
catgggtacc gaaatttett attgatggtg aatcaettga gagaatagta cgatggatga 2820
tgtatgtaga aatgacacgc aatgcaggtg ctattatcca ggaagacgac tttggaccta 2940
                                                                2960
agccaaaatt ttttatatga
```

```
<210> 158
```

## <400> 158

```
Met Ser Ser Leu Asp Glu Asp Glu Glu Asp Phe Glu Met Leu Asp Thr
1 5 10 15
```

```
Glu Asn Leu Gln Phe Met Gly Lys Lys Met Phe Gly Lys Gln Ala Gly
20 25 30
```

Glu Asp Glu Ser Asp Asp Phe Ala Ile Gly Gly Ser Thr Pro Thr Asn

<sup>&</sup>lt;211> 819

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Saccharomyces cerevisiae

40

- Lys Leu Lys Phe Tyr Pro Tyr Ser Asn Asn Lys Leu Thr Arg Ser Thr 50 55 60
- Gly Thr Leu Asn Leu Ser Leu Ser Asn Thr Ala Leu Ser Glu Ala Asn 65 70 75 80
- Gly Lys Asp Glu Glu Ser Val Asp Ser Arg Ile Lys Arg Trp Ser Pro 100 105 110
- Phe His Glu Asn Glu Ser Val Thr Thr Pro Ile Thr Lys Arg Ser Ala 115 120 125
- Glu Lys Thr Asn Ser Pro Ile Ser Leu Lys Gln Trp Asn Gln Arg Trp 130 135 140
- Phe Pro Lys Asn Asp Ala Arg Thr Glu Asn Thr Ser Ser Ser Ser Ser 145 150 155 160
- Tyr Ser Val Ala Lys Pro Asn Gln Ser Ala Phe Thr Ser Ser Gly Leu 165 170 175
- Val Ser Lys Met Ser Met Asp Thr Ser Leu Tyr Pro Ala Lys Leu Arg 180 185 190
- Ile Pro Glu Thr Pro Val Lys Lys Ser Pro Leu Val Glu Gly Arg Asp 195 200 205
- His Lys His Val His Leu Ser Ser Ser Lys Asn Ala Ser Ser Ser Leu 210 215 220
- Ser Val Ser Pro Leu Asn Phe Val Glu Asp Asn Asn Leu Gln Glu Asp 225 230 235 240
- Leu Leu Phe Ser Asp Ser Pro Ser Ser Lys Ala Leu Pro Ser Ile His 245 250 255
- Val Pro Thr Ile Asp Ser Ser Pro Leu Ser Glu Ala Lys Tyr His Ala 260 265 270
- His Asp Arg His Asn Asn Gln Thr Asn Ile Leu Ser Pro Thr Asn Ser 275 280 285
- Leu Val Thr Asn Ser Ser Pro Gln Thr Leu His Ser Asn Lys Phe Lys

295

300

Lys Ile Lys Arg Ala Arg Asn Ser Val Ile Leu Lys Asn Arg Glu Leu 305 310 315 320

Thr Asn Ser Leu Gln Gln Phe Lys Asp Asp Leu Tyr Gly Thr Asp Glu 325 330 335

Asn Phe Pro Pro Pro Ile Ile Ile Ser Ser His His Ser Thr Arg Lys 340 345 350

Asn Pro Gln Pro Tyr Gln Phe Arg Gly Arg Tyr Asp Asn Asp Thr Asp 355 360 365

Glu Glu Ile Ser Thr Pro Thr Arg Arg Lys Ser Ile Ile Gly Ala Thr 370 375 380

Ser Gln Thr His Arg Glu Ser Arg Pro Leu Ser Leu Ser Ser Ala Ile 385 390 395 400

Val Thr Asn Thr Thr Ser Ala Glu Thr His Ser Ile Ser Ser Thr Asp 405 410 415

Ser Ser Pro Leu Asn Ser Lys Arg Leu Ile Ser Ser Asn Lys Leu 420 425 430

Ser Ala Asn Pro Asp Ser His Leu Phe Glu Lys Phe Thr Asn Val His 435 440 445

Ser Ile Gly Lys Gly Gln Phe Ser Thr Val Tyr Gln Val Thr Phe Ala 450 455 460

Gln Thr Asn Lys Lys Tyr Ala Ile Lys Ala Ile Lys Pro Asn Lys Tyr 465 470 475 480

Asn Ser Leu Lys Arg Ile Leu Leu Glu Ile Lys Ile Leu Asn Glu Val 485 490 495

Thr Asn Gln Ile Thr Met Asp Gln Glu Gly Lys Glu Tyr Ile Ile Asp 500 505 510

Tyr Ile Ser Ser Trp Lys Phe Gln Asn Ser Tyr Tyr Ile Met Thr Glu 515 520 525

Leu Cys Glu Asn Gly Asn Leu Asp Gly Phe Leu Gln Glu Gln Val Ile 530 535 540

Ala Lys Lys Lys Arg Leu Glu Asp Trp Arg Ile Trp Lys Ile Ile Val

545					550					555				!	560
Glu :	Leu	Ser	Leu	Ala 565	Leu	Arg	Phe	Ile	His 570	Asp	Ser	Cys :	His :	Ile ' 575	Val
His	Leu	Asp	Leu 580	Lys	Pro	Ala	Asn	Val 585	Met	Ile	Thr	Phe	Glu ( 590	Gly .	Asn
Leu	Lys	Leu 595	Gly	Asp	Phe	Gly	Met 600	Ala	Thr	His	Leu	Pro 605	Leu	Glu	Asp
Lys	Ser 610	Phe	Glu	Asn	Glu	Gly 615	Asp	Arg	Glu	Tyr	Ile 620	Ala	Pro	Glu	Ile
Ile 625	Ser	Asp	Cys	Thr	Tyr 630	Asp	Tyr	Lys	Ala	Asp 635	Ile	Phe	Ser	Leu	Gly 640
Leu	Met	Ile	Val	Glu 645	Ile	Ala	Ala	Asn	Val 650	Val	Leu	Pro	Asp	Asn 655	Gly
Asn	Ala	Trp	His		Leu	Arg	Ser	Gly 665		Leu	Ser	Asp	Ala 670	Gly	Arg
Leu	Ser	Ser 675		Asp	Ile	His	Ser 680		Ser	Leu	Phe	Ser 685	Asp	Ile	Thr
Lys	Val		Thr	: Asn	Asp	Leu 695		. Asp	Phe	e Glu	700	Asp	Asn	Ile	Ser
Gly 705		. Sei	r Ası	n Asr	710		Thr	: Sei	Thi	r Val	L His	Asn	Asn	Ser	720
Ile	Asr	ı Ası	n Pro	725		: Asr	n Asr	ı Gly	y Ası 73	n Ası	) Asn	Asr	Asn	Val 735	Asn
Thr	· Ala	a Al	a Thi		s Ası	a Arg	g Lei	1 Ile 74		u Hi	s Lys	s Sei	750	Lys )	: Ile
Pro	Ala	a Tr		l Pr	o Ly:	s Pho	e Le	u Il O	e As	p Gl	y Glı	3 Set 76!	r Leu	ı Glu	a Arg
Ile	• Va		g Tr	р Ме	t Il	e Gl		o As	n Ty	r Gl	u Arg 78	g Ar	g Pro	Th:	r Ala
Ası	n Gl	n Il	e Le	u Gl	n Th	r Gl	u Gl	u Cy	s Le	u Ty	r Va	1 G1	u Me	t Th	r Arg

Asn Ala Gly Ala Ile Ile Gln Glu Asp Asp Phe Gly Pro Lys

785

Printed:17-08-2000

790

815

Phe Phe Ile

<210> 159

<211> 809

<212> DNA

<213> Saccharomyces cerevisiae

<400> 159

aattcccgca tgggtaccga aatttcttat tgatggtgaa tcacttgaga gaatagtacg 60 atggatgata gagcccaatt atgagagaag gcccacggca aatcaaatct tacaaactga 120 ggaatgcctg tatgtagaaa tgacacgcaa tgcaggtgct attatccagg aagacgactt 180 tggacctaag ccaaaatttt ttatatgata aatggaacaa aaaaccttgt tttatttaca 240 tactttttc ccacacgtgc ttatgggccg cattgtataa ataatccaat aacgaaaaag 300 agtgtaattg cagtccggta gtaataccat gtaaaacctt agatgagttt attttaagta 360 cagccgcttc aagcatttt attttattt tacagatgta gcagataaca accgttaaat 420 tatattatat atatatata atatataca aatacgacgt attacatata tattgagaat 480 aagggaagga tggaagacaa atgacaaaaa gtttgaagca taaatatgtt cttcgcttag 540 atgttcaatc tggttcttct ccagtttctt ctcttagcgt tgtaacggat aggtttgttg 600 gttctcaatc tgaacgact ttgagcctaa ttaaaaggga aacatatcgt gcacatacga 720 aggttcaaat tgtaaaaaat gttagtaaca atgttcaacc tcatcaatat gatgcattca 780 cggatccaag gcaataccac ctgacataa

<210> 160

<211> 102

<212> PRT

<213> Saccharomyces cerevisiae

<400> 160

Met Thr Lys Ser Leu Lys His Lys Tyr Val Leu Arg Leu Asp Val His

1 5 10 15

Leu Gly Ser Ser Pro Val Ser Ser Leu Ser Val Val Thr Asp Ser Val
20 25 30

Val Gly Ser Gln Ser Asp Pro Leu Trp Gln Trp Ser Val Leu Leu Leu 35 40 45

Ser Leu Ser His Phe Leu Leu Asp Ser Glu Arg Leu Leu Ser Leu Ile 50 55 60

Lys Arg Glu Thr Tyr Arg Ala His Thr Lys Cys Thr Ile Val Lys Asn

SEQL

65

70

75

80

Val Ser Asn Asn Val Gln Thr His Gln Tyr Asp Ala Phe Thr Asp Pro 85 90 95

Arg Gln Tyr His Leu Thr 100

<210> 161

<211> 1042

<212> DNA

<213> Saccharomyces cerevisiae

#### <400> 161

tattcaagaa ttattcaca tcttccagcg agtaaacatg ccgctggtaa tcgcgcgtcc 60 tcaatatttg agtttttcaa atagtgaggt gtggatgtat agaggaatta cacactttta 120 agtatgtgat gtatgggcgc acagtaccaa tttaactttt ttttttttc attttttagc 180 ttgattttca aaaaacttat gggcgtttta ggcteegget caaactacca ccaccacgeg 240 gcaggccgag gcaaacagta cgccttggcg gggacgccga agcgactcct tctgttccaa 300 geteaatggt eettgegttt aegetegege gtgggetaae taaegeaatt eggettttgg 360 gctgtcgaga accgagaatt attcttcgcc ttgatagata ctttaaaact tctacttaat 420 atactttcta caatttttgg tacattcata ttatactgaa aattcgaaaa agacaagcaa 480 ataaacacag atagatcaac atggctgtat gttagaaaga tattataaat cccagttaga 540 tgctgaactg atcaatagca aattataaac cacatccatc taaatgacct taccacctac 600 aatttggatt tgaaatagaa gcaatgtgta aaatataggg aaaggattag gagtgttaac 660 catactaaaa tttttcttat ccgaaacaga aatctaaagt cgccactacg cagattaaaa 720 tatggtcata aactgcttat tctgagaact tttggtggtc cagcgtggtt tatgtcaggt 780 ggtattgcct tggatccgtg aatgcatcat attgatgagt ttgaacattg ttactaacat 840 tttttacaat tgtacacttc gtatgtgcac gatatgtttc ccttttaatt aggctcaaaa 900 gtctttcaga atcaagcaaa aaatggctaa ggctaagaag caaaacagac cattgccaca 960 atggatcaga ttgagaacca acaacactat ccgttacaac gctaagagaa gaaactggag 1020 aagaaccaag atgaacatct aa 1042

<210> 162

<211> 51

<212> PRT

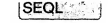
<213> Saccharomyces cerevisiae

<400> 162

Met Ala Ala Gln Lys Ser Phe Arg Ile Lys Gln Lys Met Ala Lys Ala 1 5 10 15

Lys Lys Gln Asn Arg Pro Leu Pro Gln Trp Ile Arg Leu Arg Thr Asn
20 25 30

5 1 1 2 . 5 . 1 Zal



```
Asn Thr Ile Arg Tyr Asn Ala Lys Arg Arg Asn Trp Arg Arg Thr Lys
35 40 45
```

Met Asn Ile 50

<210> 163

<211> 893

<212> DNA

<213> Saccharomyces cerevisiae

<400> 163

tctcgacagc ccaaaagccg aattgcgtta gttagcccac gcgcgagcgt aaacgcaagg 60 accattgagc ttggaacaga aggagtcgct tcggcgtcc cgccaaggcg tactgtttgc 120 ctcggcctgc cgcgtggtgg tggtagtttg agccggagcc taaaacgccc ataagtttt 180 tgaaaatcaa gctaaaaaat gaaaaaaaaa aaaagttaaa ttggtactgt gcgcccatac 240 atcacatact taaaagtgtg taattcctct atacatccac acctcactat ttgaaaaact 300 caaatattga ggacgcgga ttaccagcgg catgtttact cgctggaaga tgtgaaataa 360 ttcttgaata tgggtttgca gttagtcaat ttcaatctaa ataatcttaa tcgactattc 420 aattcttaaa ttgtaaggtt ttttaatggc ctttcagttc tagtatttt tataaaacaa 480 gaccaacata catatccaag atgaccagat cttccgtttt agctgatgct ttgaatgcca 540 ttacaagct tttgaaagcc ggtaagcgtc aagtttaat cagaccatcc tccaaggtca 600 ttatcaagtt tttgcaagtt atgcaaaagc acggttacat tggtgaattt gaatacatcg 660 atgaccacag atctggtaag attgttgtc aattgaacgg tagattgaac aagtgtggtg 720 ttatttcccc aagattcaac gttaagattg gtgacattga aaaatggact gccaacttgt 780 tgccagccag acaattcggt tacgtcatct tgaccacctc tgctggtatc atggaccatg 840 aagaagccag aagaaagcac gtttctggta agattttagg tttcgtttac taa 893

<210> 164

<211> 130

<212> PRT

<213> Saccharomyces cerevisiae

<400> 164

Met Thr Arg Ser Ser Val Leu Ala Asp Ala Leu Asn Ala Ile Asn Asn 1 5 10 15

Ala Glu Lys Thr Gly Lys Arg Gln Val Leu Ile Arg Pro Ser Ser Lys
20 25 30

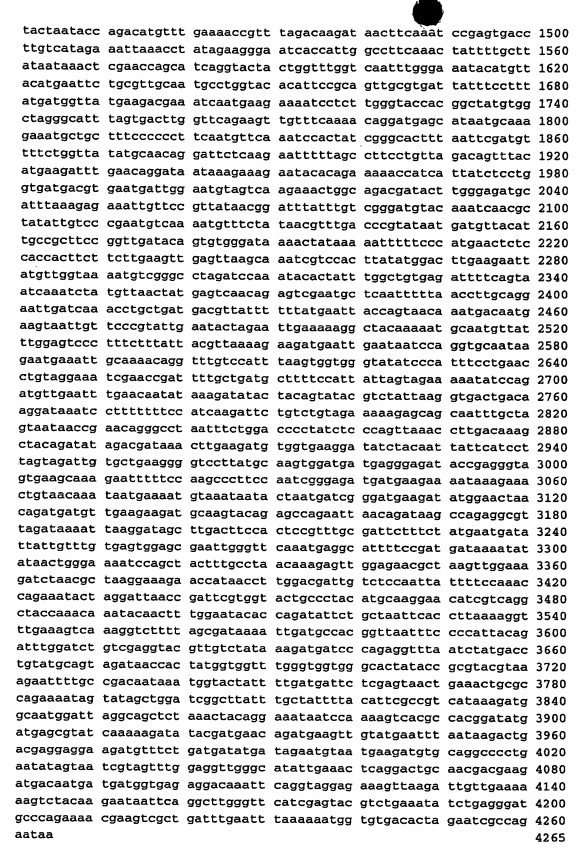
Val Ile Ile Lys Phe Leu Gln Val Met Gln Lys His Gly Tyr Ile Gly 35 40 45

Glu Phe Glu Tyr Ile Asp Asp His Arg Ser Gly Lys Ile Val Val Gln
50 55 60

```
Leu Asn Gly Arg Leu Asn Lys Cys Gly Val Ile Ser Pro Arg Phe Asn
                     70
                                          75
                                                              80
65
Val Lys Ile Gly Asp Ile Glu Lys Trp Thr Ala Asn Leu Leu Pro Ala
                                                          95
                 85
                                      90
Arg Gln Phe Gly Tyr Val Ile Leu Thr Thr Ser Ala Gly Ile Met Asp
            100
                                 105
                                                     110
His Glu Glu Ala Arg Arg Lys His Val Ser Gly Lys Ile Leu Gly Phe
                            120
        115
Val Tyr
    130
<210> 165
<211> 4265
<212> DNA
<213> Saccharomyces cerevisiae
<400> 165
cttcttcggg gttccccaat gtcactgcca tctctaaagt agatagaaag ggcgatcgct 60
```

atttaagtgt catgactttc ttaacgagag gtgttccagc ttctattttg gctttcctat 120 gtgtcattac tctaggttat ggtatcatgg catctgttgt caagggtaac gcaacctctg 180 cgtaagagat acgtctacga tagcatgtaa cagcgtatat ataatctata tgtattcgat 240 taatccaaca cttctgttgt gtagtctaat gttcgaaaaa aaggtacgct tccctgttat 300 aatcaggtat atttcgttat tcttataagc taaaagatta aaaatttttc cactttcctt 360 gaaatttggt cggttcgtgg aaaaatatta ttacgtattg aagaagtgca tgaagataaa 420 agatgggatt actggaaaaa taaagggagg aaaatcctgc agaacgttgt tgtttcaatc 480 gaaggtttct tcattcgaaa atgggttctt cagatgtttc aagtcgtgaa tgttcattgg 540 tttataatga agatcccgat ttcaccgatg gcacaacacc atgtgatcgg ttgggagtgg 600 acttgatgaa tgttctagat gacaaggatg aaataaagca agagtctgtc ccagtctcag 660 atcgtgaaat tgaggatacg gaatccgatg cttccgctgt ttcttcattt gctagcgcta 720 atgagttaat agctgagcca cacgctgcaa gtgaaactaa tcttggaact aatggtcaag 780 atggaagaaa tgtcttagaa caacaaagag acgtggttgc tagactaata gaagaaaaca 840 aggaaacgca aaaagagggt gataaagtct gtattgtccc caaggtttgg tacgataaat 900 ttttcgaccc cgatgttacc gatcctgaag atataggccc tattaataca cgcatgattt 960 gcagagactt tgaaaatttt gtgcttgagg attacaatag atgtccgtat ctgtctattg 1020 cagageetgt tttcaatttt etateagaaa tttaeggeat gacaagtgge tettaeceag 1080 tggtaactaa tttggttatc aaccaaacca caggggagtt agagacagaa tacaataaat 1140 ggtttttcag actacattac ctgaccgaaa aacaagacgg gaggaaaagg aggcatggcc 1200 aggacgattc aataatgtac ctctcgatgt ctgcgttgaa tttagtacgt gatttggttg 1260 aaaagagtat gaatctgttt tttgagaaag ctgatcatct agacgtgaat gcggtggatt 1320 ttaaaatttg gtttgtgtca gagggatctg atattgccac agatagcaat gttagtactt 1380 ttttgaattc ttcatatgaa ataactccgc ttcaatttct cgaactaccg ataaagaaac 1440

SEQL



Printed:17-08-2000 202

<210>	166
-------	-----

<211> 1254

<212> PRT

<213> Saccharomyces cerevisiae

<400> 166

Met Gly Ser Ser Asp Val Ser Ser Arg Glu Cys Ser Leu Val Tyr Asn 1 5 10 15

Glu Asp Pro Asp Phe Thr Asp Gly Thr Thr Pro Cys Asp Arg Leu Gly 20 25 30

Val Asp Leu Met Asn Val Leu Asp Asp Lys Asp Glu Ile Lys Gln Glu 35 40 45

Ser Val Pro Val Ser Asp Arg Glu Ile Glu Asp Thr Glu Ser Asp Ala
50 55 60

Ser Ala Val Ser Ser Phe Ala Ser Ala Asn Glu Leu Ile Ala Glu Pro 65 70 75 80

His Ala Ala Ser Glu Thr Asn Leu Gly Thr Asn Gly Gln Asp Gly Arg 85 90 95

Asn Val Leu Glu Gln Gln Arg Asp Val Val Ala Arg Leu Ile Glu Glu 100 105 110

Asn Lys Glu Thr Gln Lys Glu Gly Asp Lys Val Cys Ile Val Pro Lys 115 120 125

Val Trp Tyr Asp Lys Phe Phe Asp Pro Asp Val Thr Asp Pro Glu Asp 130 135 140

Ile Gly Pro Ile Asn Thr Arg Met Ile Cys Arg Asp Phe Glu Asn Phe 145 150 155 160

Val Leu Glu Asp Tyr Asn Arg Cys Pro Tyr Leu Ser Ile Ala Glu Pro 165 170 175

Val Phe Asn Phe Leu Ser Glu Ile Tyr Gly Met Thr Ser Gly Ser Tyr 180 185 190

Pro Val Val Thr Asn Leu Val Ile Asn Gln Thr Thr Gly Glu Leu Glu 195 200 205

Thr Glu Tyr Asn Lys Trp Phe Phe Arg Leu His Tyr Leu Thr Glu Lys

ישר או ואחת

210 215 220

Gln Asp Gly Arg Lys Arg Arg His Gly Gln Asp Asp Ser Ile Met Tyr 225 230 235 240

Leu Ser Met Ser Ala Leu Asn Leu Val Arg Asp Leu Val Glu Lys Ser 245 250 255

Met Asn Leu Phe Phe Glu Lys Ala Asp His Leu Asp Val Asn Ala Val 260 265 270

Asp Phe Lys Ile Trp Phe Val Ser Glu Gly Ser Asp Ile Ala Thr Asp 275 280 285

Ser Asn Val Ser Thr Phe Leu Asn Ser Ser Tyr Glu Ile Thr Pro Leu 290 295 300

Gln Phe Leu Glu Leu Pro Ile Lys Lys Leu Leu Ile Pro Asp Met Phe 305 310 315 320

Glu Asn Arg Leu Asp Lys Ile Thr Ser Asn Pro Ser Asp Leu Val Ile 325 330 335

Glu Ile Lys Pro Ile Glu Gly Asn His His Trp Pro Ser Asn Tyr Phe 340 345 350

Ala Tyr Asn Lys Leu Glu Pro Ala Ser Gly Thr Thr Gly Leu Val Asn 355 360 365

Leu Gly Asn Thr Cys Tyr Met Asn Ser Ala Leu Gln Cys Leu Val His 370 375 380

Ile Pro Gln Leu Arg Asp Tyr Phe Leu Tyr Asp Gly Tyr Glu Asp Glu 385 390 395 400

Ile Asn Glu Glu Asn Pro Leu Gly Tyr His Gly Tyr Val Ala Arg Ala
405
410
415

Phe Ser Asp Leu Val Gln Lys Leu Phe Gln Asn Arg Met Ser Ile Met 420 425 430

Gln Arg Asn Ala Ala Phe Pro Pro Ser Met Phe Lys Ser Thr Ile Gly
435 440 445

His Phe Asn Ser Met Phe Ser Gly Tyr Met Gln Gln Asp Ser Gln Glu 450 455 460

Phe Leu Ala Phe Leu Leu Asp Ser Leu His Glu Asp Leu Asn Arg Ile

480 475 470 465 Ile Lys Lys Glu Tyr Thr Glu Lys Pro Ser Leu Ser Pro Gly Asp Asp 490 485 Val Asn Asp Trp Asn Val Val Lys Lys Leu Ala Asp Asp Thr Trp Glu 510 505 500 Met His Leu Lys Arg Asn Cys Ser Val Ile Thr Asp Leu Phe Val Gly 520 515 Met Tyr Lys Ser Thr Leu Tyr Cys Pro Glu Cys Gln Asn Val Ser Ile 535 530 Thr Phe Asp Pro Tyr Asn Asp Val Thr Leu Pro Leu Pro Val Asp Thr 555 550 545 Val Trp Asp Lys Thr Ile Lys Ile Phe Pro Met Asn Ser Pro Pro Leu 570 Leu Leu Glu Val Glu Leu Ser Lys Ser Ser Thr Tyr Met Asp Leu Lys 585 580 Asn Tyr Val Gly Lys Met Ser Gly Leu Asp Pro Asn Thr Leu Phe Gly 605 600 595 Cys Glu Ile Phe Ser Asn Gln Ile Tyr Val Asn Tyr Glu Ser Thr Glu 615 610 Ser Asn Ala Gln Phe Leu Thr Leu Gln Glu Leu Ile Lys Pro Ala Asp 630 625 Asp Val Ile Phe Tyr Glu Leu Pro Val Thr Asn Asp Asn Glu Val Ile 650 645 Val Pro Val Leu Asn Thr Arg Ile Glu Lys Gly Tyr Lys Asn Ala Met 660

Leu Phe Gly Val Pro Phe Phe Ile Thr Leu Lys Glu Asp Glu Leu Asn 675 680 685

Asn Pro Gly Ala Ile Arg Met Lys Leu Gln Asn Arg Phe Val His Leu 690 695 700

Ser Gly Gly Tyr Ile Pro Phe Pro Glu Pro Val Gly Asn Arg Thr Asp 705 710 715 720

Phe Ala Asp Ala Phe Pro Leu Leu Val Glu Lys Tyr Pro Asp Val Glu

SEQL

725	730	735

Phe	Glu	Gln	Tyr	Lys	Asp	Ile	Leu	Gln	Tyr	Thr	Ser	Ile	Lys	Val	Thr
			740					745					750		

- Asp Lys Asp Lys Ser Phe Phe Ser Ile Lys Ile Leu Ser Val Glu Lys
  755 760 765
- Glu Gln Gln Phe Ala Ser Asn Asn Arg Thr Gly Pro Asn Phe Trp Thr 770 775 780
- Pro Ile Ser Gln Leu Asn Leu Asp Lys Ala Thr Asp Ile Asp Asp Lys 785 790 795 800
- Leu Glu Asp Val Val Lys Asp Ile Tyr Asn Tyr Ser Ser Leu Val Asp 805 810 815
- Cys Ala Glu Gly Val Leu Met Gln Val Asp Asp Glu Gly Asp Thr Glu 820 825 830
- Gly Ser Glu Ala Lys Asn Phe Ser Lys Pro Phe Gln Ser Gly Asp Asp 835 840 845
- Glu Glu Asn Lys Glu Thr Val Thr Asn Asn Glu Asn Val Asn Asn Thr 850 855 860
- Asn Asp Arg Asp Glu Asp Met Glu Leu Thr Asp Asp Val Glu Glu Asp 865 870 875 880
- Ala Ser Thr Glu Pro Glu Leu Thr Asp Lys Pro Glu Ala Leu Asp Lys 885 890 895
- Ile Lys Asp Ser Leu Thr Ser Thr Pro Phe Ala Ile Leu Ser Met Asn 900 905 910
- Asp Ile Ile Val Cys Glu Trp Ser Glu Leu Gly Ser Asn Glu Ala Phe 915 920 925
- Ser Asp Asp Lys Ile Tyr Asn Trp Glu Asn Pro Ala Thr Leu Pro Asn 930 935 940
- Lys Glu Leu Glu Asn Ala Lys Leu Glu Arg Ser Asn Ala Lys Glu Arg 945 950 955 960
- Thr Ile Thr Leu Asp Asp Cys Leu Gln Leu Phe Ser Lys Pro Glu Ile 965 970 975
- Leu Gly Leu Thr Asp Ser Trp Tyr Cys Pro Thr Cys Lys Glu His Arg

980

985

- Gln Ala Thr Lys Gln Ile Gln Leu Trp Asn Thr Pro Asp Ile Leu Leu 995 1000 1005
- Ile His Leu Lys Arg Phe Glu Ser Gln Arg Ser Phe Ser Asp Lys Ile 1010 1015 1020
- Asp Ala Thr Val Asn Phe Pro Ile Thr Asp Leu Asp Leu Ser Arg Tyr 1025 1030 1035 1040
- Val Val Tyr Lys Asp Asp Pro Arg Gly Leu Ile Tyr Asp Leu Tyr Ala 1045 1050 1055
- Val Asp Asn His Tyr Gly Gly Leu Gly Gly Gly His Tyr Thr Ala Tyr 1060 1065 1070
- Val Lys Asn Phe Ala Asp Asn Lys Trp Tyr Tyr Phe Asp Asp Ser Arg 1075 1080 1085
- Val Thr Glu Thr Ala Pro Glu Asn Ser Ile Ala Gly Ser Ala Tyr Leu 1090 1095 1100
- Leu Phe Tyr Ile Arg Arg His Lys Asp Gly Asn Gly Leu Gly Ser Ser 1105 1110 1115 1120
- Lys Leu Gln Glu Ile Ile Gln Lys Ser Arg His Gly Tyr Asp Glu Arg 1125 1130 1135
- Ile Lys Lys Ile Tyr Asp Glu Gln Met Lys Leu Tyr Glu Phe Asn Lys 1140 1145 1150
- Thr Asp Glu Glu Glu Asp Val Ser Asp Asp Met Ile Glu Cys Asn Glu 1155 1160 1165
- Asp Val Gln Ala Pro Glu Tyr Ser Asn Arg Ser Leu Glu Val Gly His 1170 1175 1180
- Ile Glu Thr Gln Asp Cys Asn Asp Glu Asp Asp Asn Asp Gly Glu 1185 1190 1195 1200
- Arg Thr Asn Ser Gly Arg Arg Lys Leu Arg Leu Leu Lys Lys Val Tyr 1205 1210 1215
- Lys Asn Asn Ser Gly Leu Gly Ser Ser Ser Thr Ser Glu Ile Ser Glu 1220 1225 1230
- Gly Cys Pro Glu Asn Glu Val Ala Asp Leu Asn Leu Lys Asn Gly Val

SEQL

1235 1240 1245

Thr Leu Glu Ser Pro Glu 1250

<210> 167 <211> 3146 <212> DNA <213> Saccharomyces cerevisiae

<400> 167

tgtcgtattt ccactgattt ggatgtatgt tgcatggtgc tagattgtga ataaatcatt 60 tccactggtt tctagcagag gtaagaaaat cagtagtagc ggcagcagcc aagaaaaatg 120 gcagcctggc aagcagcgaa ggccatctgt gctgcatttc cacactcctt gtatgactgc 180 atacgcataa agaggtgcct tgctgtaggc gtatatgatc ctaagcaaca gagaaaccac 240 ggttctcttc tcttcttatt tcgttattgt cttccctttt tactatgggt aaagtcgccc 300 taaagcgggg cgctcacaat atcgccgcag ctacagccgt ttttttttt tttgttttt 360 tttgcgctgc ttctcgaaga atgaatggct cactgaaaaa ttttgattca tcgatataaa 420 gaacactgtt cacttcgatg tcatccggcc aataaagttg tttttaggat aaacgagtaa 480 gtggtagctg gtacaggatc atgagatttt cacacttctt gaagtacaat gctgtcccag 540 aatggcaaaa ccattatatg gactacagcg agctgaaaaa tcttatttac acgctacaaa 600 cagatgaact tcaggttggt gataacgaag aaggatttgg cgcaggaaag agctctaaca 660 ttacagatag gttcaaaaac aagttttctt ttaaaaatgc gaaggaagat acgtcttccg 720 gtatgaacaa agatgcaggc atcgttgagg aaaccatcga gttgcgagag ttgcctactg 780 ctcagacggt cgctgccaaa ccttctcctt tcagaagaat gaaggaaaag atattttaca 840 aaagaaggtc gtcttccgca tcgtccgtct cctccacggc caacgaaaat ctgcaattag 900 acacttatga tacgtttgtt ggtgatttaa cagctgaaaa acagaaagta gatgattttt 960 ataagaggac agaagcgaag ttctacgaca aatttgacgc gctggtgaag gacctgaaga 1020 aaatcggagt tatagaatac gatatcgacg atgatactct gtttaacgaa ccgattgcca 1080 gcacaaatga cgaagttccc ccactagact tggatgatga cgaagacgac gacgaatttt 1140 acgatgatca atctaatatt gaagataata ctgctttgct gcatcattcg cagtataaca 1200 ttaagtctca gaaaaaatcg ctgttgaaga agtcgatcgt aaacctatat atcgatcttt 1260 gccagttgaa gtcgttcatc gaattgaacc gcattgggtt tgcaaaaatt acaaagaaat 1320 cagacaaagt tcttcacttg aatacaagaa ccgaactgat cgaatcggag cagtttttca 1380 aagacacata tgcattccag gcagaaacga tcgaattgct aaattccaaa atttcccagc 1440 tagtcacatt ttatgcgcgc atcactgacc ggcctcataa tatctcgcat agcaagcaag 1500 agttgaaatc ctacctgcat gaccacattg tttgggaaag aagtaacact tggaaagaca 1560 tgttgggact gctatcgcaa gctgacgagt tgacaccaaa ggaaacagaa tataatgcaa 1620 ataagetggt aggeaagtta gatttggaat actacagatg gecactacce agacegataa 1680 acttaaaatt cactagtata aacaacgttg cactaccgaa attattttc accaagaaag 1740 catacaagat ttactttatt attctagtca ctggactctt gttaggaata aagaccttca 1800 acgacgctgc tcagcaccgc tgcatggccc ttgtcgagtg tgtcgccttt ttgtgggcta 1860 gtgaggccat cccattacac attacagcat tccttgtacc actacttgta gtccttttca 1920 aagteetaaa aaceteegae ggggetataa tgagtgetge aagegettea teagaaattt 1980 tggccgccat gtggtcttct acaattatga ttctgctggc aggttttact ttgggtgaag 2040 tacttgcaca atataacatc gccaaagttc ttgcctcgtg gttgttggcc ttcgctggtt 2100

```
gtaaacccag aaacgttctt ttaatggcaa tgtgtgtcgt gttcttccta tcaatgtgga 2160
tttccaatgt cgcagcacct gttctaacat attcgttgtt atctccccta ttggatgcca 2220
tggatgcaga tagcccattt gcgcaagcat tggtgttagg tgtagcgttg gctgcaaata 2280
teggtggtat gtetteacea atetetteac etcaaaacat catttecatg tegtaettga 2340
aaccetatgg tattggetgg ggccaattet ttgetgttge attgccatet ggtateetgg 2400 ·
ccatgctttt ggtttggatt ttattgttca ctactttcaa gatgaataag accaaattgg 2460
aaaaatttaa gcctattaag acgaaattca cagttaagca gtattatatc attactgtca 2520
ctgtggccac tattttgttg tggtgtgtgg aaagccagat tgaaggtgct tttgggtcat 2580
caggicaaat tgcaatcatt cccatcgitt tgttttttgg taccggatta ctatcaacac 2640
aagatttaaa tgcctttccg tggtcaatcg ttattttggc aatgggaggt attgctttgg 2700
ggaaggccgt ctcatcctcg ggtttgctat caaccattgc aaaagcatta caaaagaaaa 2760
ttgagaatga tggtgttttt gccattctat gtattttcgg tatcctgatg ttggttgtgg 2820
gtactttcgt ctcgcataca gtatccgcta ttatcatcat tcccttggtg caagaagttg 2880
gtgacaagct tggcaacccc aaagctgctc ctatccttgt tttcggttgt gcattattgt 2940
catcctgtgg tatgggacta gcttcttcgg ggttccccaa tgtcactgcc atctctaaag 3000
tagatagaaa gggcgatcgc tatttaagtg tcatgacttt cttaacgaga ggtgttccag 3060
cttctatttt ggctttccta tgtgtcatta ctctaggtta tggtatcatg gcatctgttg 3120
tcaagggtaa cgcaacctct gcgtaa
                                                                  3146
```

```
<210> 168
```

<211> 881

<212> PRT

<213> Saccharomyces cerevisiae

<400> 168

Met Arg Phe Ser His Phe Leu Lys Tyr Asn Ala Val Pro Glu Trp Gln
1 5 10 15

Asn His Tyr Met Asp Tyr Ser Glu Leu Lys Asn Leu Ile Tyr Thr Leu 20 25 30

Gln Thr Asp Glu Leu Gln Val Gly Asp Asn Glu Glu Gly Phe Gly Ala 35 40 45

Gly Lys Ser Ser Asn Ile Thr Asp Arg Phe Lys Asn Lys Phe Ser Phe 50 55 60

Lys Asn Ala Lys Glu Asp Thr Ser Ser Gly Met Asn Lys Asp Ala Gly 65 70 75 80

Ile Val Glu Glu Thr Ile Glu Leu Arg Glu Leu Pro Thr Ala Gln Thr
85 90 95

Val Ala Ala Lys Pro Ser Pro Phe Arg Arg Met Lys Glu Lys Ile Phe 100 105 110

Tyr Lys Arg Arg Ser Ser Ser Ala Ser Ser Val Ser Ser Thr Ala Asn

Glu Asn Leu Gln Leu Asp Thr Tyr Asp Thr Phe Val Gly Asp Leu Thr Ala Glu Lys Gln Lys Val Asp Asp Phe Tyr Lys Arg Thr Glu Ala Lys Phe Tyr Asp Lys Phe Asp Ala Leu Val Lys Asp Leu Lys Lys Ile Gly Val Ile Glu Tyr Asp Ile Asp Asp Asp Thr Leu Phe Asn Glu Pro Ile Ala Ser Thr Asn Asp Glu Val Pro Pro Leu Asp Leu Asp Asp Asp Glu Asp Asp Asp Glu Phe Tyr Asp Asp Gln Ser Asn Ile Glu Asp Asn Thr Ala Leu Leu His His Ser Gln Tyr Asn Ile Lys Ser Gln Lys Lys Ser Leu Leu Lys Lys Ser Ile Val Asn Leu Tyr Ile Asp Leu Cys Gln Leu Lys Ser Phe Ile Glu Leu Asn Arg Ile Gly Phe Ala Lys Ile Thr Lys Lys Ser Asp Lys Val Leu His Leu Asn Thr Arg Thr Glu Leu Ile Glu Ser Glu Gln Phe Phe Lys Asp Thr Tyr Ala Phe Gln Ala Glu Thr Ile Glu Leu Leu Asn Ser Lys Ile Ser Gln Leu Val Thr Phe Tyr Ala Arg Ile Thr Asp Arg Pro His Asn Ile Ser His Ser Lys Gln Glu Leu Lys Ser Tyr Leu His Asp His Ile Val Trp Glu Arg Ser Asn Thr Trp Lys 

Asp Met Leu Gly Leu Leu Ser Gln Ala Asp Glu Leu Thr Pro Lys Glu

Thr Glu Tyr Asn Ala Asn Lys Leu Val Gly Lys Leu Asp Leu Glu Tyr

370

375

380

Tyr Arg Trp Pro Leu Pro Arg Pro Ile Asn Leu Lys Phe Thr Ser Ile 385 390 395 400

Asn Asn Val Ala Leu Pro Lys Leu Phe Phe Thr Lys Lys Ala Tyr Lys 405 410 415

Ile Tyr Phe Ile Ile Leu Val Thr Gly Leu Leu Leu Gly Ile Lys Thr
420 425 430

Phe Asn Asp Ala Ala Gln His Arg Cys Met Ala Leu Val Glu Cys Val 435 440 445

Ala Phe Leu Trp Ala Ser Glu Ala Ile Pro Leu His Ile Thr Ala Phe 450 455 460

Leu Val Pro Leu Leu Val Val Leu Phe Lys Val Leu Lys Thr Ser Asp 465 470 475 480

Gly Ala Ile Met Ser Ala Ala Ser Ala Ser Ser Glu Ile Leu Ala Ala 485 490 495

Met Trp Ser Ser Thr Ile Met Ile Leu Leu Ala Gly Phe Thr Leu Gly 500 505 510

Glu Val Leu Ala Gln Tyr Asn Ile Ala Lys Val Leu Ala Ser Trp Leu 515 520 525

Leu Ala Phe Ala Gly Cys Lys Pro Arg Asn Val Leu Leu Met Ala Met 530 535

Cys Val Val Phe Phe Leu Ser Met Trp Ile Ser Asn Val Ala Ala Pro 545 550 560

Val Leu Thr Tyr Ser Leu Leu Ser Pro Leu Leu Asp Ala Met Asp Ala 565 570 575

Asp Ser Pro Phe Ala Gln Ala Leu Val Leu Gly Val Ala Leu Ala Ala 580 585 590

Asn Ile Gly Gly Met Ser Ser Pro Ile Ser Ser Pro Gln Asn Ile Ile 595 600 605

Ser Met Ser Tyr Leu Lys Pro Tyr Gly Ile Gly Trp Gly Gln Phe Phe 610 615 620

Ala Val Ala Leu Pro Ser Gly Ile Leu Ala Met Leu Leu Val Trp Ile

SEQL

625					630					635					640
Leu	Leu	Phe	Thr	Thr 645	Phe	Lys	Met	Asn	Lys 650	Thr	Lys	Leu	Glu	Lys 655	Phe
Lys	Pro	Ile	Lys 660	Thr	Lys	Phe	Thr	Val 665	Lys	Gln	Tyr	Tyr	Ile 670	Ile	Thr
Val	Thr	Val 675	Ala	Thr	Ile	Leu	Leu 680	Trp	Сув	Val	Glu	Ser 685	Gln	Ile	Glu
Gly	Ala 690	Phe	Gly	Ser	Ser	Gly 695	Gln	Ile	Ala	Ile	Ile 700	Pro	Ile	Val	Lev
Phe 705	Phe	Gly	Thr	Gly	Leu 710	Leu	Ser	Thr	Gl'n	Asp 715	Leu	Asn	Ala	Phe	Pro 720
Trp	Ser	Ile	Val	Ile 725	Leu	Ala	Met	Gly	Gly 730	Ile	Ala	Leu	Gly	Lys 735	Ala
Val	Ser	Ser	Ser 740	Gly	Leu	Leu	Ser	Thr 745	Ile	Ala	Lys	Ala	Leu 750	Gln	Lys
Lys	Ile	Glu 755	Asn	Asp	Gly	Val	Phe 760	Ala	Ile	Leu	Cys	Ile 765	Phe	Gly	Ile
Leu	Met 770	Leu	Val	Val	Gly	Thr 775	Phe	Val	Ser	His	Thr 780	Val	Ser	Ala	Ile
Ile 785	Ile	Ile	Pro	Leu	Val 790	Gln	Glu	Val	Gly	Asp 795	Lys	Leu	Gly	Asn	Pro 800
Lys	Ala	Ala	Pro	Ile 805	Leu	Val	Phe	Gly	Cys 810	Ala	Leu	Leu	Ser	Ser 815	Суя
Gly	Met	Gly	Leu 820	Ala	Ser	Ser	Gly	Phe 825	Pro	Asn	Val	Thr	Ala 830	Ile	Se
Lys	Val	Asp 835	Arg	Lys	Gly	Asp	Arg 840	Tyr	Leu	Ser	Val	Met 845	Thr	Phe	Let
Thr	Arg 850	Gly	Val	Pro	Ala	Ser 855	Ile	Leu	Ala	Phe	Leu 860	Cys	Val	Ile	Th
Leu 865	Gly	Tyr	Gly	Ile	Met 870	Ala	Ser	Val	Val	Lys 875	Gly	Asn	Ala	Thr	Se:

Ala

<210> 169

```
<211> 2093
<212> DNA
<213> Saccharomyces cerevisiae
<400> 169
actgccatct cttccattct ttgaaaagat tattaagttc gtcgttaagt tccgcaatct 60
gcatttcatt taaagacgga tcatatattc ttgtgctttt ttgttttatc tctttactta 120
cttgtcgctt ccattcgttg gcctctttta tggattttac cttagacacg ttcctgggcc 180
tctggtaacg tgaataatcc ttgtatccac ctgccgactc tgcttgctgc tcttgaaatc 240
gtaccagaac tgagttggcc ttatctacat ttctactcat tttgaatgca caggtatctg 300
attactgatg tggtgcgcct tgcgatataa cggtgtgtca cttttatttg ctctttcatg 360
catcctgaaa ttatttcacc gcactacgca aagagaacgg agaaaaaggt ataatagggc 420
aaatgatcat tgacatcgtg atcgtaagcc aaaaaaaaat acaataggct ccctaaataa 480
gtagagtaaa agctcttgag atgaaggaga atgacatgaa taatggcgta gataaatggg 540
taaatgagga agatggtcga aatgatcatc ataacaacaa taataacttg atgaagaagg 600
ccatgatgaa caatgagcaa attgatagaa ctcaggatat cgacaacgcc aaagaaatgt 660
tgaggaaaat atcaagtgaa agcagctcgc gcagaagctc cctgttgaat aaagattcat 720
ctctcgtgaa cggcaatgca aacagtggcg gtggtacgag cattaacgga acaagaggaa 780
gttctaagag tagtaataca cactttcagt atgcctccac ggcgtatggt gtaagaatgt 840
tgagtaaaga tatatctaat accaaagtgg aactggatgt ggaaaatttg atgattgtta 900
cgaaactcaa cgatgtctca ctgtatttct taacaagaga gttggtagaa tgggttttgg 960
tacattttcc acgtgtgact gtttatgtgg attccgaatt gaaaaacagc aaaaaatttg 1020
ccgctggcga gttatgtgaa gatagtaaat gtagagaatc aaggatcaag tattggacaa 1080
aggatttcat cagggaacat gatgttttct tcgatttggt agtgactttg ggtggcgacg 1140
gtactgttct ttttgtaagt tccatttttc agagacatgt accacccgtt atgtcgtttt 1200
cattagggtc tctaggattt ttaacaaatt ttaagtttga acatttcagg gaggatttac 1260
ctcggattat gaatcataaa atcaagacaa atttacggtt gaggttggag tgcacaattt 1320
atcgtagaca ccgccctgaa gtagacccaa acacggggaa gaaaatatgt gtggtggaaa 1380
aactaagcac acaccacatt ttgaacgaag tgaccatcga tcgtggtcca agtccttttc 1440
tatccatgtt agaattgtat ggtgacggct cattaatgac cgttgcgcag gcggacggac 1500
tgattgctgc tactccgact gggtccacgg cctattcttt gagtgcaggt gggtcattgg 1560
tatgcccaac cgtcaatgca atcgctttaa cacccatttg tccacatgca ttgagtttca 1620
gacccatcat cttaccagaa agtataaatt taaaagtgaa agtctcgatg aagtcaaggg 1680
ctccagcatg ggcggctttt gatgggaaag atagaattga attgcaaaaa ggtgatttta 1740
taaccatatg cgccagccca tatgcttttc caaccgtgga agcctcgccc gatgagttta 1800
ttaacagtat cagtcgacaa ctaaactgga atgtgaggga acaacaaaag tcctttacgc 1860
atattttgtc ccaaaagaac caagaaaaat atgcacatga ggcgaacaaa gtcagaaatc 1920
aagcagaacc tttagaggta ataagagata aatactctct ggaagcagac gctactaagg 1980
aaaacaacaa cggaagcgat gatgagagcg acgatgagag tgtaaactgc gaagcttgca 2040
aattaaagcc ttcgagcgtc ccaaaacctt ctcaagcaag gttttcagta taa
                                                                  2093
```

<210> 170

<211> 530

<212> PRT

<213> Saccharomyces cerevisiae

<400> 170

Met Lys Glu Asn Asp Met Asn Asn Gly Val Asp Lys Trp Val Asn Glu
1 5 10 15

Glu Asp Gly Arg Asn Asp His His Asn Asn Asn Asn Asn Leu Met Lys
20 25 30

Lys Ala Met Met Asn Asn Glu Gln Ile Asp Arg Thr Gln Asp Ile Asp 35 40 45

Asn Ala Lys Glu Met Leu Arg Lys Ile Ser Ser Glu Ser Ser Ser Arg
50 55 60

Arg Ser Ser Leu Leu Asn Lys Asp Ser Ser Leu Val Asn Gly Asn Ala 65 70 75 80

Asn Ser Gly Gly Gly Thr Ser Ile Asn Gly Thr Arg Gly Ser Ser Lys 85 90 95

Ser Ser Asn Thr His Phe Gln Tyr Ala Ser Thr Ala Tyr Gly Val Arg 100 105 110

Met Leu Ser Lys Asp Ile Ser Asn Thr Lys Val Glu Leu Asp Val Glu 115 120 125

Asn Leu Met Ile Val Thr Lys Leu Asn Asp Val Ser Leu Tyr Phe Leu 130 135 140

Thr Arg Glu Leu Val Glu Trp Val Leu Val His Phe Pro Arg Val Thr 145 150 155 160

Val Tyr Val Asp Ser Glu Leu Lys Asn Ser Lys Lys Phe Ala Ala Gly 165 170 175

Glu Leu Cys Glu Asp Ser Lys Cys Arg Glu Ser Arg Ile Lys Tyr Trp 180 185 190

Thr Lys Asp Phe Ile Arg Glu His Asp Val Phe Phe Asp Leu Val Val
195 200 205

Thr Leu Gly Gly Asp Gly Thr Val Leu Phe Val Ser Ser Ile Phe Gln 210 215 220

Arg His Val Pro Pro Val Met Ser Phe Ser Leu Gly Ser Leu Gly Phe

225	230	235	240
220			

Leu Thr Asn Phe Lys Phe Glu His Phe Arg Glu Asp Leu Pro Arg Ile 245 250 255

Met Asn His Lys Ile Lys Thr Asn Leu Arg Leu Arg Leu Glu Cys Thr 260 265 270

Ile Tyr Arg Arg His Arg Pro Glu Val Asp Pro Asn Thr Gly Lys Lys 275 280 285

Ile Cys Val Val Glu Lys Leu Ser Thr His His Ile Leu Asn Glu Val 290 295 300

Thr Ile Asp Arg Gly Pro Ser Pro Phe Leu Ser Met Leu Glu Leu Tyr 305 310 315 320

Gly Asp Gly Ser Leu Met Thr Val Ala Gln Ala Asp Gly Leu Ile Ala 325 330 335

Ala Thr Pro Thr Gly Ser Thr Ala Tyr Ser Leu Ser Ala Gly Gly Ser 340

Leu Val Cys Pro Thr Val Asn Ala Ile Ala Leu Thr Pro Ile Cys Pro 355 360 365

His Ala Leu Ser Phe Arg Pro Ile Ile Leu Pro Glu Ser Ile Asn Leu 370 375 380

Lys Val Lys Val Ser Met Lys Ser Arg Ala Pro Ala Trp Ala Ala Phe 385 390 395 400

Asp Gly Lys Asp Arg Ile Glu Leu Gln Lys Gly Asp Phe Ile Thr Ile 405 410 415

Cys Ala Ser Pro Tyr Ala Phe Pro Thr Val Glu Ala Ser Pro Asp Glu
420 425 430

Phe Ile Asn Ser Ile Ser Arg Gln Leu Asn Trp Asn Val Arg Glu Gln 435

Gln Lys Ser Phe Thr His Ile Leu Ser Gln Lys Asn Gln Glu Lys Tyr 450 455 460

Ala His Glu Ala Asn Lys Val Arg Asn Gln Ala Glu Pro Leu Glu Val 465 470 475 480

Ile Arg Asp Lys Tyr Ser Leu Glu Ala Asp Ala Thr Lys Glu Asn Asn

```
485 490 495
```

```
Asn Gly Ser Asp Asp Glu Ser Asp Glu Ser Val Asn Cys Glu Ala
500 505 510
```

Cys Lys Leu Lys Pro Ser Ser Val Pro Lys Pro Ser Gln Ala Arg Phe 515 520 525

Ser Val

```
<210> 171
<211> 1255
<212> DNA
<213> Saccharomyces cerevisiae
```

# <400> 171

```
aggttgtgtt cttcaatgat gggcaatgca atttggcgtt aagcgcctga gcaataaggt 60
aacagcgaaa tttatgacat attatttcga accttttaca aactagtaga tttagtgatt 120
tattacctat tggcattcat ttgtgttcta tatgtggatg aggatagccg cctttcttct 180
categgagge catateatet ttegacaate etttttaaat aetattteea teegtgeete 240
taatagattt gtgtagttgt ctgggtgcaa tctttccatt tttgctgaac ttttttttt 300
ttttcatgtt tttcagattc tgaagtaccg caataggata tggcggataa tccgccatat 360
gatecgeete atactageea ttacecatet ateceaggea ttatgggtat geaacteata 420
atctcaaata cacaaataag agcaacctta tatatcactt tttcccgttc agcaagaggt 480
aaagccacca aaggttcaaa atgcaaatgt atgttacggc gaatacagaa tactatgttc 540
gaaataatat gaggattata cgatagcaaa aaagccataa acgaaagaca taaatggaaa 600
atgattgaca agctcacaat ttattaaaca agtagcaatt gagaaaaact attactcgcg 660
gcaagcttct gagtttacat taaatctgta gagcaaattg aaaatgtcgc atatgtgctg 720
aagggtttgt ttgttccatc ttattttgca taacatagtt atatttactt ggtcgcataa 780
aaaatatttt ttactaacgt gaagtttctt tctttatgat gtacgcacgc acgtctgtgc 840
ttactccata aatgaactta ttccaatttt gtacagcttc gttaagactt tgactggtaa 900
gaccatcact ttggaagttg aatcttctga caccattgac aatgtcaagt ccaagatcca 960
agacaaggaa ggtatcccac ctgaccaaca aagattgatc tttgctggta agcaattgga 1020
agacggtaga actctatctg actacaacat ccaaaaggaa tccactttac atttggtctt 1080
gagattaaga ggtggtatca ttgaaccatc tttgaaagcc ttggcttcca aatacaactg 1140
tgacaaatct gtttgtcgta aatgttacgc cagattacca ccaagagcta ccaactgtag 1200
aaagagaaag tgtggtcaca ccaaccaatt gcgtccaaag aagaagttga aataa
```

```
<210> 172
<211> 128
<212> PRT
<213> Saccharomyces cerevisiae
```

<400> 172

```
Met Gln Ile Phe Val Lys Thr Leu Thr Gly Lys Thr Ile Thr Leu Glu
                                                          15
                                     10
  1
Val Glu Ser Ser Asp Thr Ile Asp Asn Val Lys Ser Lys Ile Gln Asp
             20
                                 25
Lys Glu Gly Ile Pro Pro Asp Gln Gln Arg Leu Ile Phe Ala Gly Lys
                             40
Gln Leu Glu Asp Gly Arg Thr Leu Ser Asp Tyr Asn Ile Gln Lys Glu
                         55
Ser Thr Leu His Leu Val Leu Arg Leu Arg Gly Gly Ile Ile Glu Pro
                                         75
                     70
Ser Leu Lys Ala Leu Ala Ser Lys Tyr Asn Cys Asp Lys Ser Val Cys
                                                          95
Arg Lys Cys Tyr Ala Arg Leu Pro Pro Arg Ala Thr Asn Cys Arg Lys
            100
                                105
                                                    110
Arg Lys Cys Gly His Thr Asn Gln Leu Arg Pro Lys Lys Leu Lys
        115
                            120
                                                125
```

```
<210> 173
<211> 1175
<212> DNA
<213> Saccharomyces cerevisiae
```

# <400> 173

```
ageteteaaa caactaatae tataagttea agtacaagea caggaggtgt aggtteagte 60
aagccatgtc tttacttcgt tttaatgtta gaaacaatcg cttatttgtt ttcttaaaca 120
aatatattag gttcaaggtc ttcgcaggtg taagaaaacc cgtggtctcc atattcttaa 180
gtatgataaa taaaaaaaaa cttaataaat tattaattgc ttcaaacctt tttcttttt 240
tagtttttaa tatttcaaac gttatcttca ttgaacgccc aaatagggaa aaatcctggc 300
aaatttttta ttgctgtcat ccaaggctat gctagaaaat tcaagagctt ggatgattta 360
aaaagacact ctcaatcgag aaagtttatt ctttgttatt ctgctttacc tgatcatatt 420
ccggcgtatt gtttctaatc aagtgatttc gatatccagt tacgaaccat ttacaacatt 480
cctgaaaata ttgcgtatca atgatatttg ctccttcttt ctccctcatt aaaaatattc 540
tcctggtaag ctttctaatc agccacagtt ttgctgccaa aactttaacg tctagttcca 600
atgacgatac acttgccagg tccgcagctg cagatgcaga catggcattc ttcatggagt 660
ttttaaacga tttcgacacc gcttttccac agtatacctc atacatgatg caaaaccatt 720
taaccctacc tcaacctgtt gctgactact actatcacat ggttgatttg gcctcaacag 780
```

217 Printed:17-08-2000

1175

cagatttaca atctgatatt gctcagagtt ttccgttcac tcaattccaa acattcatta 840 eggeetttee atggtatace tetttgetaa acaaageete egeeaceace atatacette 900 cccaacactt cataacaggt gagacagaag ctaccatgac taactcatct tatgccagcc 960 aaaaaaactc cgtttccaat tctgttcctt tctcgacagc gaacgcaggc cagtccatga 1020 tttccatggc taatgaagaa aacagtacaa cagcacttat atccgcatca aactcttctt 1080 caacatccag aactagtcaa tcacagaatg gtgcccatgc caaaagctta tatttcccca 1140 tggcgttgtt cggaatcttt gcagttgccc tttaa <210> 174 <211> 224 <212> PRT <213> Saccharomyces cerevisiae <400> 174 Met Ile Phe Ala Pro Ser Phe Ser Leu Ile Lys Asn Ile Leu Leu Val 10 Ser Phe Leu Ile Ser His Ser Phe Ala Ala Lys Thr Leu Thr Ser Ser 20 25 Ser Asn Asp Asp Thr Leu Ala Arg Ser Ala Ala Ala Asp Ala Asp Met 40 Ala Phe Phe Met Glu Phe Leu Asn Asp Phe Asp Thr Ala Phe Pro Gln 50 Tyr Thr Ser Tyr Met Met Gln Asn His Leu Thr Leu Pro Gln Pro Val 65 70 75 Ala Asp Tyr Tyr His Met Val Asp Leu Ala Ser Thr Ala Asp Leu 85 90 Gln Ser Asp Ile Ala Gln Ser Phe Pro Phe Thr Gln Phe Gln Thr Phe 100 105

Ile Thr Ala Phe Pro Trp Tyr Thr Ser Leu Leu Asn Lys Ala Ser Ala 120

Thr Thr Ile Tyr Leu Pro Gln His Phe Ile Thr Gly Glu Thr Glu Ala 130 135 140

Thr Met Thr Asn Ser Ser Tyr Ala Ser Gln Lys Asn Ser Val Ser Asn 145 150 160

Ser Val Pro Phe Ser Thr Ala Asn Ala Gly Gln Ser Met Ile Ser Met 165 170 175

```
Ala Asn Glu Glu Asn Ser Thr Thr Ala Leu Ile Ser Ala Ser Asn Ser
180 185 190
```

Ser Ser Thr Ser Arg Thr Ser Gln Ser Gln Asn Gly Ala His Ala Lys 195 . 200 205

Ser Leu Tyr Phe Pro Met Ala Leu Phe Gly Ile Phe Ala Val Ala Leu 210 215 220

<210> 175 <211> 1618 <212> DNA <213> Saccharomyces cerevisiae

#### <400> 175

gcaccctttt tcttttcaa taacaaaaaa gagcttgggc attacgtttt caacactttt 60 aatatttccc tttttacatc cgtacatata aaatgtaaac ccatgcattc atgacaattt 120 tacctattte ttaaaggeet ttgtcaette tttatgggaa tggegetaet ttaattttee 180 gcctactttg aaaaatttac caaggcgaaa ttgagtgcgc taggcggaag ttccagggca 240 cggtcaccga acctttgtgc tgtttcgaac gagggtcccg ctggaggttg acggacgcgg 300 gaggagctgg aaagatggaa tggaaggact gcaacactca gtaagaaggt tcgtggtagg 360 gaacatcacc ttctctagtt cctgtaaaaa taatatgatc aagtttgaat cctcgtattg 420 aaaaatcgcg agttattaag tgtgtaattt agaataccga aatagcacaa gaagagataa 480 gataagatac tgataggata atgtctttac cagctacttt tgatttgact ccagaggatg 540 cccaactttt gttggccgct aacacccatt taggtgctag aaacgttcaa gtatgtacac 600 atatcccata cgattatgtt ctatagatga taataggtct cgaaaagaat atgtccccga 660 tttaatcata tttggaggtc agaggaccaa aggtaatttc agaggaattt tgaacacgcc 720 gggtttagaa gagttagaat ttcacttctt agtgagggta aggagaagaa actcaataag 780 aatatacatc ctcgactgtg tcaaagatta tgaactccga tgaaacagta aaacgtcaaa 840 aaattccacg ggatataatt cggtacttta ttggatatgg aagaaaacat tatatatgca 900 caaatgaacg ctcttactaa cataatttat ctttcctctc tttttttagg tccaccaaga 960 accatacgtt ttcaatgcta gaccagatgg tgttcacgtt atcaatgttg gtaagacctg 1020 ggaaaagttg gttttggctg ctagaattat tgctgccatt ccaaacccag aagatgttgt 1080 tgccatctct tccagaacct acggtcaaag agctgtcttg aaatttgctg ctcacactgg 1140 tgctactcca atcgccggta gattcactcc aggttctttc actaattaca tcacccgttc 1200 tttcaaggaa ccaagattgg ttattgttac cgacccaaga ttagacgctc aggccattaa 1260 ggaagettet taegttaaca ttecagteat tgetttgaet gatttggaet ceceatetga 1320 atttgttgat gtcgccatcc catgtaacaa cagaggtaag cactccatcg gtttaatctg 1380 gtacttgttg gctagagaag ttttgagact aagaggtgct ttggtcgaca gaactcaacc 1440 atggtccatc atgccagatt tgtacttcta cagaaaccca gaagaagttg agcaagttgc 1500 tgaagaagct gctgccgctg aagaaggtga agaagaagaa gttaaggaag aagtcactga 1560 aggtcaagct gaagctactg aatgggctga agaaaatgca gacaacgttg aatggtaa

<210> 176

<211> 252

<212> PRT

<213> Saccharomyces cerevisiae

<400> 176

Met Ser Leu Pro Ala Thr Phe Asp Leu Thr Pro Glu Asp Ala Gln Leu
1 5 10 15

Leu Leu Ala Ala Asn Thr His Leu Gly Ala Arg Asn Val Gln Val His
20 25 30

Gln Glu Pro Tyr Val Phe Asn Ala Arg Pro Asp Gly Val His Val Ile 35 40 45

Asn Val Gly Lys Thr Trp Glu Lys Leu Val Leu Ala Ala Arg Ile Ile 50 55 60

Ala Ala Ile Pro Asn Pro Glu Asp Val Val Ala Ile Ser Ser Arg Thr 65 70 75 80

Tyr Gly Gln Arg Ala Val Leu Lys Phe Ala Ala His Thr Gly Ala Thr
85 90 95

Pro Ile Ala Gly Arg Phe Thr Pro Gly Ser Phe Thr Asn Tyr Ile Thr
100 105 110

Arg Ser Phe Lys Glu Pro Arg Leu Val Ile Val Thr Asp Pro Arg Leu 115 120 125

Asp Ala Gln Ala Ile Lys Glu Ala Ser Tyr Val Asn Ile Pro Val Ile 130 135 140

Pro Cys Asn Asn Arg Gly Lys His Ser Ile Gly Leu Ile Trp Tyr Leu 165 170 175

Leu Ala Arg Glu Val Leu Arg Leu Arg Gly Ala Leu Val Asp Arg Thr
180 185 190

Gln Pro Trp Ser Ile Met Pro Asp Leu Tyr Phe Tyr Arg Asn Pro Glu 195 200 205

Glu Val Glu Gln Val Ala Glu Glu Ala Ala Ala Glu Glu Gly Glu 210 215 220

```
Glu Glu Glu Val Lys Glu Glu Val Thr Glu Gly Gln Ala Glu Ala Thr
225 230 235 240
```

Glu Trp Ala Glu Glu Asn Ala Asp Asn Val Glu Trp
245 250

<210> 177 <211> 2345 <212> DNA <213> Saccharomyces cerevisiae

<400> 177

cttcccatgt agaatatcat tgtcaaacat actactaccg tgagaatcca atctactaaa 60 aagacccagc taaagtcctt ttggtgtgtc aatggaacac cacgtaattg tgaaatagct 120 tccatgcttg ctaacttatt accgtcaata gaagacttga gtcaggtggc tttaatgagt 180 actattettt tttttttcc aaagagcact atgttgataa taccgcagta attttttttg 240 agtateetgt ageetagaaa ggttgaaget tataaaaceg tgeeaacage tttatagtgg 300 ggagtttggc ttccctctat ttgtatattg atcgccatcc ctacgaagtt attgggaacg 360 catcgtgaac ctctcacttt aatgccagcg gtagaaaaaa aagtcataaa caatacacgc 420 eggetacact tagaaagaaa tgacatttgc tgtcttataa aaggacttga cagaccaaag 480 acgcgtataa tacaccaaga atggccttat tggagaagtt gcatcgaagg attgttgata 540 tggggcttgt cccgcgtata atcgccttat taccagttat ttccatgcta tgcgctctat 600 ttgggtttat ttctatagct attctgccta tggatggaca gtacagaaga acatacattt 660 ctgagaatgc attgatgcct tcacaagcgt atagttactt tagagaatct gaatggaaca 720 ttttgagggg ctatcgatct caaattaaag aaatggtaaa catgacttct atggaaagaa 780 acaatttgat gggttcttgg ttacaagaat ttggtactaa gactgctatt tacgaaaatg 840 aacaatatgg agaaacattg tacggtgtaa tgcacgctcc taggggtgat ggaacagaag 900 cgatggtgct tgccgttcca tggtttaatt cagatgatga attcaatatt ggcggcgcag 960 ctttgggtgt atctttagca agatttttct cacgttggcc agtatggtcc aagaatataa 1020 ttgttgtctt cagcgaaaat cctcgtgcag cattaagatc atgggttgag gcataccata 1080 cttccttaga tttgactggt ggttccattg aagctgctgt tgtgttggat tattcgagta 1140 cggaagattt cttcgagtat gtagaaatct catacgacgg tctgaatggt gagctgccca 1200 atttggatct tgtcaacatc gctatatcca ttacggaaca tgaaggtatg aaagtttctt 1260 tgcacggtct acccagtgat cagttaacta ataataattt ctggtcaaga ttaaaaatat 1320 tatgcctggg aataagggat tgggcgttgt ccggtgttaa aaagccccat ggtaacgagg 1380 catttagcgg ctggaggatt caatctgtaa cattgaaagc acatggaaac agtggtcatg 1440 atattactac atttggacgt atacccgaag caatgtttcg ctctattaat aaccttttgg 1500 aaaaatttca ccaatcgttc ttcttttatt tgttattagc accacgtcag ttcgtatcca 1560 ttagtagtta tttgccaagc gctgtggctt tatctatagc attcgccata agttcattaa 1620 atgcatttat aaacaatgct tatgcaaata tatccttatt ttccgagtat aatttggtag 1680 cgttgttggt ttggttcgtg tcattggtga tatcatttgt tgtttcacaa gcgtttcttc 1740 taataccttc atcgggatta ttgatgacaa ttagcatggc atcttgtttt ttacctttga 1800 tactttccag aaaaattcac atctcagaac cactatcata caggttgaaa aatgttgctt 1860 ttttatattt cagtttggtt tcaacatctt tgctaatgat aaactttgca atggctttac 1920 tgatcggcac attggcattt cctatgacat ttgtgaagac cattgttgaa agttctagcg 1980

Printed:17-08-2000 221

aacatgaggt gacaactcaa teetetaace caataaaaac tgageegaaa gatgagatag 2040 agetegtega gaatcacatg gatacaacge cageaaceee ecaacaacag aaacaaaaac 2100 taaaaaattt agtactatta attttgacaa ateeatttat tteaataace ttatteggae 2160 tattttttga tgatgaattt catggatttg atataataaa caaactggtt teageatggt 2220 tggatttgaa atgttggagt tggtttgtae tttgtatagg ttggetteea tgttggetat 2280 tgatattage gteategtt gaatctaaat etgtegtagt aaggtegaaa gaaaageaaa 2340 gttag

<210> 178

<211> 614

<212> PRT

<213> Saccharomyces cerevisiae

<400> 178

Met Ala Leu Leu Glu Lys Leu His Arg Arg Ile Val Asp Met Gly Leu
1 5 10 15

Val Pro Arg Ile Ile Ala Leu Leu Pro Val Ile Ser Met Leu Cys Ala 20 25 30

Leu Phe Gly Phe Ile Ser Ile Ala Ile Leu Pro Met Asp Gly Gln Tyr 35 40 45

Arg Arg Thr Tyr Ile Ser Glu Asn Ala Leu Met Pro Ser Gln Ala Tyr 50 55 60

Ser Tyr Phe Arg Glu Ser Glu Trp Asn Ile Leu Arg Gly Tyr Arg Ser 65 70 75 80

Gln Ile Lys Glu Met Val Asn Met Thr Ser Met Glu Arg Asn Asn Leu 85 90 95

Met Gly Ser Trp Leu Gln Glu Phe Gly Thr Lys Thr Ala Ile Tyr Glu
100 105 110

Asn Glu Gln Tyr Gly Glu Thr Leu Tyr Gly Val Met His Ala Pro Arg 115 120 125

Gly Asp Gly Thr Glu Ala Met Val Leu Ala Val Pro Trp Phe Asn Ser 130 135 140

Asp Asp Glu Phe Asn Ile Gly Gly Ala Ala Leu Gly Val Ser Leu Ala 145 150 155 160

Arg Phe Phe Ser Arg Trp Pro Val Trp Ser Lys Asn Ile Ile Val Val
165 170 175

Phe	Ser	Glu	Asn	Pro	Arg	Ala	Ala	Leu	Arg	Ser	$\mathtt{Trp}$	Val	Glu	Ala	Tyr
			180					185					190		

His Thr Ser Leu Asp Leu Thr Gly Gly Ser Ile Glu Ala Ala Val Val 195 200 205

Leu Asp Tyr Ser Ser Thr Glu Asp Phe Phe Glu Tyr Val Glu Ile Ser 210 215 220

Tyr Asp Gly Leu Asn Gly Glu Leu Pro Asn Leu Asp Leu Val Asn Ile 225 230 235 240

Ala Ile Ser Ile Thr Glu His Glu Gly Met Lys Val Ser Leu His Gly 245 250 255

Leu Pro Ser Asp Gln Leu Thr Asn Asn Asn Phe Trp Ser Arg Leu Lys 260 265 270

Ile Leu Cys Leu Gly Ile Arg Asp Trp Ala Leu Ser Gly Val Lys Lys 275 280 285

Pro His Gly Asn Glu Ala Phe Ser Gly Trp Arg Ile Gln Ser Val Thr 290 295 300

Leu Lys Ala His Gly Asn Ser Gly His Asp Ile Thr Thr Phe Gly Arg 305 310 315 320

Ile Pro Glu Ala Met Phe Arg Ser Ile Asn Asn Leu Leu Glu Lys Phe 325 330 335

His Gln Ser Phe Phe Phe Tyr Leu Leu Leu Ala Pro Arg Gln Phe Val 340 345 350

Ser Ile Ser Ser Tyr Leu Pro Ser Ala Val Ala Leu Ser Ile Ala Phe 355 360 365

Ala Ile Ser Ser Leu Asn Ala Phe Ile Asn Asn Ala Tyr Ala Asn Ile 370 375 380

Ser Leu Phe Ser Glu Tyr Asn Leu Val Ala Leu Leu Val Trp Phe Val 385 390 395 400

Ser Leu Val Ile Ser Phe Val Val Ser Gln Ala Phe Leu Leu Ile Pro 405 410 415

Ser Ser Gly Leu Leu Met Thr Ile Ser Met Ala Ser Cys Phe Leu Pro 420 425 430 Leu Ile Leu Ser Arg Lys Ile His Ile Ser Glu Pro Leu Ser Tyr Arg 435 440 445

Leu Lys Asn Val Ala Phe Leu Tyr Phe Ser Leu Val Ser Thr Ser Leu 450 455 460

Leu Met Ile Asn Phe Ala Met Ala Leu Leu Ile Gly Thr Leu Ala Phe 465 470 475 480

Pro Met Thr Phe Val Lys Thr Ile Val Glu Ser Ser Ser Glu His Glu
485 490 495

Val Thr Thr Gln Ser Ser Asn Pro Ile Lys Thr Glu Pro Lys Asp Glu 500 505 510

Ile Glu Leu Val Glu Asn His Met Asp Thr Thr Pro Ala Thr Pro Gln
515 520 525

Gln Gln Lys Gln Lys Leu Lys Asn Leu Val Leu Leu Ile Leu Thr Asn 530 535 540

Pro Phe Ile Ser Ile Thr Leu Phe Gly Leu Phe Phe Asp Asp Glu Phe 545 550 555 560

His Gly Phe Asp Ile Ile Asn Lys Leu Val Ser Ala Trp Leu Asp Leu 565 570 575

Lys Cys Trp Ser Trp Phe Val Leu Cys Ile Gly Trp Leu Pro Cys Trp 580 585 590

Leu Leu Ile Leu Ala Ser Ser Phe Glu Ser Lys Ser Val Val Arg 595 600 605

Ser Lys Glu Lys Gln Ser 610

<210> 179

<211> 845

<212> DNA

<213> Saccharomyces cerevisiae

<400> 179

tccgtcccaa agatagcttg atcatcctaa gcgggtcgca ttttactatc cccagttccg 60 aagaataaca aggctcaatg catatgctgt ttatcgtgct cacaaggcgg tatgacgttt 120 ttagcaatgt aattatata aaaagatctt cagtcactat gagtggttga ttgccccata 180 gagagctata agccgacgtg aaagctgctg gttccagctt ggctcatgtc gtcaccagtc 240



```
actagteact tggtegeatt cattgetact catetgegag tgageatatt tgagatetga 300 cttgeeaagg gattagaate acgtaagaet ettgateett agaagatatt tetgaeaaag 360 aaceacetaa geeatgeaag tttttttt catttggtgg egaaacaaag gtgatgaaag 420 tttettettg tacaaaegee aageeegata ggtgagaeaa ttettgaagt aatggaeete 480 tgagtaagea tateagtega atgaagttee aatatgegtt ggeeaaggaa cagetaggea 540 geaactegeg eagtggegte aaaaaaetaa taagtaaaea eeactggett eeggaataet 600 atttetetga teteteatt tetgttgae ageagtggga eagtagagee attgaaaaaa 660 etacaaateat atettgtatg eggeeegeaa aceaagagat ttateetttg agacattgtg 720 agaeeeeteeg ttegeaaeeg tgetetetgt ttteateaet atatgeaege tettteeaaa 780 geteetgeae tttgeaegtg geggageeat egeeeggett eeatatgtae ggetgeeaea 840 eetaa
```

<210> 180

<211> 114

<212> PRT

<213> Saccharomyces cerevisiae

<400> 180

Met Lys Phe Gln Tyr Ala Leu Ala Lys Glu Gln Leu Gly Ser Asn Ser

1 5 10 15

Arg Ser Gly Val Lys Lys Leu Ile Ser Lys His His Trp Leu Pro Glu 20 25 30

Tyr Tyr Phe Ser Asp Leu Ser Phe Ser Val Val Gln Gln Trp Asp Ser

35 40 45

Arg Ala Ile Glu Lys Thr Thr Ile Ile Ser Cys Met Arg Pro Ala Asn 50 55 60

Gln Glu Ile Tyr Pro Leu Arg His Cys Glu Thr Leu Arg Ser Gln Pro 65 70 75 80

Cys Ser Leu Phe Ser Ser Leu Tyr Ala Arg Ser Phe Gln Ser Ser Cys 85 90 95

Thr Leu His Val Ala Glu Pro Ser Pro Gly Phe His Met Tyr Gly Cys
100 105 110

His Thr

<210> 181

<211> 959

<212> DNA

<213> Saccharomyces cerevisiae

```
<400> 181
 atcaaaagag tgtttcaagt aagtaaaaac atttgagcct ccccatttgt tgaaaggaga 60
 gaaattaaac ttggttgggg ttaattattt gatgggtata ttaatttgca accgcaaggt 120
 atcgataata aatattctac aaaaccttta tcaatagtgg tgaagtcttt agtgcgatct 180
 acctggggtt aatgaacgag aagttcttga gatatctttc ctgtttacct ccgtgcatcc 240
 tgtaaggaat tgggtttatc atttatcatt tattttagta caaacttttt tttttggccc 300
 gggcgcactt tttcaagcgg tgggaactca tcaaaatgaa aaactagata cttttagact 360
 tattaaatgg tttaaatatt ttgagatgtt cgttatatca gaaacttcct tacttctatc 420
 ttttattcca atacaaagaa gtcacaagat tacttggtaa gaaagaagca gttaattttt 480
 aattttgccg acaagccaag atgcaaattt tcgtcaagac tttaaccggt aagactatta 540
ccctggaagt tgaatcttct gacactattg acaatgtcaa gtccaagatc caagacaagg 600
aaggtattcc acctgaccaa caaagattga tctttgctgg taagcaattg gaagatggta 660
gaactttgtc cgactacaac atccaaaagg aatctactct acacttggtc ttgagattga 720
gaggtggtgg taagaagaga aagaagaagg tctacaccac cccaaagaag atcaagcaca 780
agcacaagaa ggtcaagttg gctgtcttgt cctactacaa ggtcgatgct gaaggtaagg 840
ttaccaaatt gagaagagaa tgtagcaacc caacttgtgg tgctggtgtt ttcttggcta 900
accacaagga cagattgtac tgtggtaagt gtcattccgt ctacaaggtt aacgcttaa 959
<210> 182
<211> 152
<212> PRT
<213> Saccharomyces cerevisiae
<400> 182
Met Gln Ile Phe Val Lys Thr Leu Thr Gly Lys Thr Ile Thr Leu Glu
  1
                  5
                                     10
                                                         15
Val Glu Ser Ser Asp Thr Ile Asp Asn Val Lys Ser Lys Ile Gln Asp
             20
                                 25
Lys Glu Gly Ile Pro Pro Asp Gln Gln Arg Leu Ile Phe Ala Gly Lys
                             40
Gln Leu Glu Asp Gly Arg Thr Leu Ser Asp Tyr Asn Ile Gln Lys Glu
                         55
Ser Thr Leu His Leu Val Leu Arg Leu Arg Gly Gly Lys Lys Arg
 65
                     70
                                         75
Lys Lys Lys Val Tyr Thr Thr Pro Lys Lys Ile Lys His Lys
                 85
                                     90
                                                         95
Lys Val Lys Leu Ala Val Leu Ser Tyr Tyr Lys Val Asp Ala Glu Gly
```

105

110

```
Lys Val Thr Lys Leu Arg Arg Glu Cys Ser Asn Pro Thr Cys Gly Ala
        115
                            120
                                                125
Gly Val Phe Leu Ala Asn His Lys Asp Arg Leu Tyr Cys Gly Lys Cys
                        135
                                            140
His Ser Val Tyr Lys Val Asn Ala
145
                    150
<210> 183
<211> 848
<212> DNA
<213> Saccharomyces cerevisiae
<400> 183
agccaccaat tccaccaggc ccggcattca agtacttgta agaacaccaa caggcaaagt 60
caacacccca atcatgaagt tgcaatggga cgttccctac agcgtgtgct aaatcccatc 120
caaccaatat gtcggggaat tggtgggcaa atgaggtgat tcggccaata tcgaaatatt 180
gccccgtgta atactgaaca cctgacaaac agaccagcgc caattcatct tgatttacct 240
ctatggtate caggatatet tgagttetga tataagtete teeetegegt ggetegatet 300
qaataaaaac attctcaggt tccgaaattc catgaatttt gcactggttg tagaaagcat 360
aatagtctga tggaaaggag cccttttcaa aaaggatctt gaatcttttc tcagtaggtt 420
tataaaacgt aattaacaat gaattcaaat ttgcagtgag actattcatt actgcaactt 480
cattttcttg agcacccacg atgggggcta gtagtggaag aataggtaag tcaatgctga 540
cccaaggcac ctttcctctg gcttcttcag gatgtttgaa atgcgattcc acagcacaat 600
cgctccacgc atctagctca gcattaattg aattcctagt tgacttcggc atcaaaccta 660
aagaattccc gcataagtac gtcacaggct tatcgtcgga cgatagtccc atggatttaa 720
atgtagggat gttgaattca tccctcagag attccggata ttctccgtct aattccaaag 780
ctttctccat aattaccagt ttgtttcttt tttccacaga atatttaggc ttaatatgta 840
                                                                   848
cgtattga
<210> 184
<211> 115
<212> PRT
<213> Saccharomyces cerevisiae
<400> 184
Met Gly Ala Ser Ser Gly Arg Ile Gly Lys Ser Met Leu Thr Gln Gly
 1
                  5
                                                          15
Thr Phe Pro Leu Ala Ser Ser Gly Cys Leu Lys Cys Asp Ser Thr Ala
             20
                                 25
                                                     30
Gln Ser Leu His Ala Ser Ser Ser Ala Leu Ile Glu Phe Leu Val Asp
```

227

40

45

SEQL

```
Phe Gly Ile Lys Pro Lys Glu Phe Pro His Lys Tyr Val Thr Gly Leu
     50
                         55
                                             60
Ser Ser Asp Asp Ser Pro Met Asp Leu Asn Val Gly Met Leu Asn Ser
                     70
                                         75
Ser Leu Arg Asp Ser Gly Tyr Ser Pro Ser Asn Ser Lys Ala Phe Ser
                 85
                                     90
Ile Ile Thr Ser Leu Phe Leu Phe Ser Thr Glu Tyr Leu Gly Leu Ile
            100
                                105
Cys Thr Tyr
        115
<210> 185
<211> 2600
<212> DNA
<213> Saccharomyces cerevisiae
<400> 185
gcgcattttc cccacccatg cgaatcatgg ctgaaatcaa tgcctgttag gtgccctgca 60
acactagtca tggtaacttc gcagttagcc ccatttctgg caaacgggaa cccactaaac 120
atgaaatcat agttctttac atacatgtag ccggaatccc ttgaagttga tctgcctcct 180
cctaggatct gtgaaactgc cttcgctata gaatttttct ctgcgacaca tagcactttc 240
attattgtct ataatacacc tttacgtagc tgaagcgagc gcacagacga aggtgctttc 300
aaaagcgatg ccctctttat tgacagttac taattcgcca aactgctttt ttttatcaat 360
gtgattttcg cgttcacgcc attatccaat gtcaaaaagg ataatggaaa tgttaacatg 420
cctgtatgat ccgaccgttt ttagcaaact tatcagggga aaaagtatat tccattaaat 480
gacacatgcc accatagata atggataatg aagaagttaa cgaagaatgt atgagattat 540
ttttcaagaa cgctcgtgcg catctggata aacatctaac atcaaggttg acatgcgatg 600
aaaatgcata tatcacgttc agatgcttcc tggatggaat acatcgcaaa tctactaggt 660
ttctcgaaga gctacttttg aaacaagaaa atatgtacca taataacaat tacgaacgca 720
taaatgattc cgtgatacca ttggttctga aacttttatg gcttcaaatt cacgaaccta 780
cactccaatg gtttgagcac tggttccatg atatcatgcg actaagtaac agaagaaagt 840
tcagagtttt tagaattttt caaaaaaaaa tgattcaatt tttcaaaatt acacacaggt 900
attactatga catcatcgaa cacctatgcg caaagtacga tatgaattcc gttatttcaa 960
atgctctctt cgcgaagttg aatttaatgc aatacacaga tggactttca actcatgaga 1020
aaattatett aaacacgagt aatecaetga egttttecat tgtaatetea etacaaagat 1080
gcgtgattaa tctaggttcc acacattttt ataaaacact actaaacaag ccgtctaaca 1140
aacccaagag tgtggaaggt tttgagaagt ctattaggta cttgaatatt gcctcactct 1200
atctcccagc cgttggagat acttattttc aacgagcgaa aatttacttg atcactggga 1260
aattctcact gtatttcttt gaattagtaa gaggagcatt ggtaaggatt ccgtctaaat 1320
```

gtgcgttaaa caatttgaaa gatttcattt tgactcctga ttttccggaa agaagacgtc 1380 tgatgaaaaa attggcaatt cttgtgtcaa aagatctcaa aggtgagaaa tcattctttg 1440

```
aaggtcaaat tgttttgcaa tttctatcga tagtagaaca cactttggtt ccacagtcat 1500
ggaacgcatc acgtgcttct aattgttggt tattgaaaga gcatttacaa atggctgcat 1560
taaagtatca ttcaggtaat attaatgtta tacttgaaaa cttggctgcc acaatgggaa 1620
gtttcgatct tatgtttaca actcgaaaaa gtaaggaaca aaagaacaaa ctcaaatatg 1680
cagatttgag tgagcgccag gtttttttt tagacttgag ctttgatttt attgctaata 1740
tcatagacgt cgtcatcaaa ccctcctggc aaaaaaacat ggaagacttt cgatatctag 1800
ccattattcg tttgcttatg tgctggatta agtcatatag atctattttg cagtacactc 1860
acagacacag gaagttttgc acttcattcg ccttgttgct gaacgacttg ataaatagtc 1920
cactgaattg ttcaggaaat atatatagcc acaggccgaa aagaagctat ctttttagag 1980
aagatattat tttcagggaa ttttcttgca ttaactttgc actaacagat tttaatgacg 2040
attatgtgta tgattetece gacatgatta ataatataat tggatgeeet acattgaeta 2100
aagtgettte tecaaaagaa gaatgtgtte tgegaattag atcaataata ttttetggea 2160
tgaaattttt agagaaaaat gacaecggcg tcatatggaa tgccagcaaa tataagtttg 2220
atttaataag cccaaatatt aaaataaaac gccaaatagc attatcggaa atttcctcca 2280
aaataaatgt aaaaacacaa caggaaagag tagtctcttc gagaaaagtt gaggccaaaa 2340
gagatgaaca acagcgcaaa agagccggga aaatagctgt gacagaactg gaaaaacaat 2400
ttgcaaatgt ccggagaaca aaaaaattgt ctccgctccc agaaaaagat ggcgtttctt 2460
ctgagttggt aaaacatgct gcttcacgag ggagaaaaac tatcactggc ccactatcct 2520
ctgattttct ctcatatcca gacgaagcaa ttgatgctga tgaggacatc accgtccaag 2580
                                                                  2600
tgccagatac tcctacttga
<210> 186
<211> 699
<212> PRT
<213> Saccharomyces cerevisiae
```

<400> 186

Met Asp Asn Glu Glu Val Asn Glu Glu Cys Met Arg Leu Phe Phe Lys

1 5 10 15

Asn Ala Arg Ala His Leu Asp Lys His Leu Thr Ser Arg Leu Thr Cys
20 25 30

Asp Glu Asn Ala Tyr Ile Thr Phe Arg Cys Phe Leu Asp Gly Ile His 35 40 45

Arg Lys Ser Thr Arg Phe Leu Glu Glu Leu Leu Leu Lys Gln Glu Asn 50 55 60

Met Tyr His Asn Asn Asn Tyr Glu Arg Ile Asn Asp Ser Val Ile Pro 65 70 75 80

Leu Val Leu Lys Leu Leu Trp Leu Gln Ile His Glu Pro Thr Leu Gln 85 90 95

Trp Phe Glu His Trp Phe His Asp Ile Met Arg Leu Ser Asn Arg Arg 100 105 110 Lys Phe Arg Val Phe Arg Ile Phe Gln Lys Lys Met Ile Gln Phe Phe 115 120 125

Lys Ile Thr His Arg Tyr Tyr Tyr Asp Ile Ile Glu His Leu Cys Ala 130 135 140

Lys Tyr Asp Met Asn Ser Val Ile Ser Asn Ala Leu Phe Ala Lys Leu 145 150 155 160

Asn Leu Met Gln Tyr Thr Asp Gly Leu Ser Thr His Glu Lys Ile Ile 165 170 175

Leu Asn Thr Ser Asn Pro Leu Thr Phe Ser Ile Val Ile Ser Leu Gln
180 185 190

Arg Cys Val Ile Asn Leu Gly Ser Thr His Phe Tyr Lys Thr Leu Leu 195 200 205

Asn Lys Pro Ser Asn Lys Pro Lys Ser Val Glu Gly Phe Glu Lys Ser 210 225 220

Ile Arg Tyr Leu Asn Ile Ala Ser Leu Tyr Leu Pro Ala Val Gly Asp 225 230 235 240

Thr Tyr Phe Gln Arg Ala Lys Ile Tyr Leu Ile Thr Gly Lys Phe Ser 245 250 255

Leu Tyr Phe Phe Glu Leu Val Arg Gly Ala Leu Val Arg Ile Pro Ser 260 265 270

Lys Cys Ala Leu Asn Asn Leu Lys Asp Phe Ile Leu Thr Pro Asp Phe 275 280 285

Pro Glu Arg Arg Leu Met Lys Lys Leu Ala Ile Leu Val Ser Lys 290 295 300

Asp Leu Lys Gly Glu Lys Ser Phe Phe Glu Gly Gln Ile Val Leu Gln 305 310 315 320

Phe Leu Ser Ile Val Glu His Thr Leu Val Pro Gln Ser Trp Asn Ala 325 330 335

Ser Arg Ala Ser Asn Cys Trp Leu Leu Lys Glu His Leu Gln Met Ala 340 345 350

Ala Leu Lys Tyr His Ser Gly Asn Ile Asn Val Ile Leu Glu Asn Leu 355 360 365



- Ala Ala Thr Met Gly Ser Phe Asp Leu Met Phe Thr Thr Arg Lys Ser 370 375 380
- Lys Glu Gln Lys Asn Lys Leu Lys Tyr Ala Asp Leu Ser Glu Arg Gln 385 390 395 400
- Val Phe Phe Leu Asp Leu Ser Phe Asp Phe Ile Ala Asn Ile Ile Asp 405 410 415
- Val Val Ile Lys Pro Ser Trp Gln Lys Asn Met Glu Asp Phe Arg Tyr 420 425 430
- Leu Ala Ile Ile Arg Leu Leu Met Cys Trp Ile Lys Ser Tyr Arg Ser 435 440 445
- Ile Leu Gln Tyr Thr His Arg His Arg Lys Phe Cys Thr Ser Phe Ala 450 455 460
- Leu Leu Asn Asp Leu Ile Asn Ser Pro Leu Asn Cys Ser Gly Asn 465 470 475 480
- Ile Tyr Ser His Arg Pro Lys Arg Ser Tyr Leu Phe Arg Glu Asp Ile 485 490 495
- The Phe Arg Glu Phe Ser Cys Ile Asn Phe Ala Leu Thr Asp Phe Asn 500 505 510
- Asp Asp Tyr Val Tyr Asp Ser Pro Asp Met Ile Asn Asn Ile Ile Gly 515 520 525
- Cys Pro Thr Leu Thr Lys Val Leu Ser Pro Lys Glu Glu Cys Val Leu 530 535 540
- Arg Ile Arg Ser Ile Ile Phe Ser Gly Met Lys Phe Leu Glu Lys Asn 545 550 555 560
- Asp Thr Gly Val Ile Trp Asn Ala Ser Lys Tyr Lys Phe Asp Leu Ile 565 570 575
- Ser Pro Asn Ile Lys Ile Lys Arg Gln Ile Ala Leu Ser Glu Ile Ser 580 585 590
- Ser Lys Ile Asn Val Lys Thr Gln Gln Glu Arg Val Val Ser Ser Arg 595 600 605
- Lys Val Glu Ala Lys Arg Asp Glu Gln Gln Arg Lys Arg Ala Gly Lys 610 615 620



```
Ile Ala Val Thr Glu Leu Glu Lys Gln Phe Ala Asn Val Arg Arg Thr 625 630 635 640
```

Lys Lys Leu Ser Pro Leu Pro Glu Lys Asp Gly Val Ser Ser Glu Leu 645 650 655

Val Lys His Ala Ala Ser Arg Gly Arg Lys Thr Ile Thr Gly Pro Leu 660 665 670

Ser Ser Asp Phe Leu Ser Tyr Pro Asp Glu Ala Ile Asp Ala Asp Glu 675 680 685

Asp Ile Thr Val Gln Val Pro Asp Thr Pro Thr 690 695

<210> 187

<211> 2471

<212> DNA

<213> Saccharomyces cerevisiae

### <400> 187

aatggtatca cggaatcatt tatgcgttcg taattgttat tatggtacat attttcttgt 60 ttcaaaagta gctcttcgag aaacctagta gatttgcgat gtattccatc caggaagcat 120 ctgaacgtga tatatgcatt ttcatcgcat gtcaaccttg atgttagatg tttatccaga 180 tgcgcacgag cgttcttgaa aaataatctc atacattctt cgttaacttc ttcattatcc 240 attatctatg gtggcatgtg tcatttaatg gaatatactt tttcccctga taagtttgct 300 aaaaacggtc ggatcataca ggcatgttaa catttccatt atcctttttg acattggata 360 atggcgtgaa cgcgaaaatc acattgataa aaaaaagcag tttggcgaat tagtaactgt 420 caataaagag ggcatcgctt ttgaaagcac cttcgtctgt gcgctcgctt cagctacgta 480 aaggtgtatt atagacaata atgaaagtgc tatgtgtcgc agagaaaaat tctatagcga 540 aggcagtttc acagatccta ggaggaggca gatcaacttc aagggattcc ggctacatgt 600 atgtaaagaa ctatgatttc atgtttagtg ggttcccgtt tgccagaaat ggggctaact 660 gcgaagttac catgactagt gttgcagggc acctaacagg cattgatttc agccatgatt 720 cgcatgggtg gggaaaatgc gccatccaag agttatttga tgcgccactg aacgagatta 780 tgaataacaa ccaaaaaaag atagcaagca acatcaagcg agaagcgagg aatgcagact 840 atctgatgat atggacagat tgcgaccggg aaggagagta catcggttgg gagatatggc 900 aggaggccaa gagaggcaac aggctcatac aaaatgatca agtataccgg gcagtctttt 960 cgcatctcga aagacaacac atattaaatg cagcacgaaa cccaagtcga ttggatatga 1020 agagtgtgca cgctgtaggc acgcggattg aaatcgatct tcgagcaggt gttacattca 1080 ccagactctt aacagaaacg ctacgaaata aactgagaaa ccaagccacc atgaccaagg 1140 atggtgcaaa acaccgcggt ggtaacaaga acgactcaca agtcgtatcg tatggtacat 1200 gccagtttcc aacgctcggc tttgtagtag acaggtttga aagaatacga aattttgttc 1260 ccgaagagtt ctggtatatc caattggtag tcgaaaacaa agacaacggc ggaacaacaa 1320 cgttccagtg ggacaggggc cacttgttcg accggctgag cgtgttaacg ttttacgaga 1380 catgcatcga aaccgccggc aatgttgctc aagtagtaga cttgaaatca aagccaacaa 1440 cgaaatacag acctttacct ctgaccacag tggagctaca aaaaaactgc gcccggtacc 1500

```
tgcgtctgaa cgccaaacaa tcactagacg cagcagaaaa gctataccaa aaggggttca 1560
tatcgtatcc aagaacagag actgatactt tcccacacgc aatggaccta aaatccttgg 1620
tcgaaaagca agctcaattg gaccaactcg ctgcaggcgg cagaaccgcc tgggcatcgt 1680
acgcggcatc gctgctccaa cccgaaaaca caagtaacaa taacaagttc aagtttccac 1740
gaagcggctc ccatgacgac aaagcgcatc caccaatcca ccccatcgta agtctggggc 1800
ctgaagcaaa tgtttcgcca gtggaaagaa gagtatacga gtacgtggcc aggcactttt 1860
tggcatgctg ctcagaggac gccaagggcc aatcgatgac ccttgtgttg gactgggccg 1920
ttgaacgttt ctcagcttca ggtctcgtag tcctagagag aaatttcctc gatgtttacc 1980
cttgggcccg atgggaaacc accaagcagt taccgcggct tgaaatgaat gccctcgtag 2040
acatcgcgaa ggccgaaatg aaggcgggca ctacggcgcc gcccaagccg atgactgaga 2100
gtgaactcat teteetcatg gatacaaacg geattggeac agaegecace attgeggage 2160
acatagacaa gatccaagta cgtaattacg ttaggagcga gaaagtaggc aaggaaacct 2220
acttacaacc cacgaccetg ggtgteteac tagtgcacgg ettegaggee ateggeeteg 2280
aagactcctt tgcaaagccc ttccagcgca gagaaatgga gcaagacctc aagaaaatct 2340
gcgaaggtca tgcctccaag actgatgttg taaaggacat agtcgagaag tataggaagt 2400
actggcacaa gacgaatgcc tgcaagaata ctctcttgca agtttatgac cgtgtcaagg 2460
                                                                   2471
catccatgta a
<210> 188
<211> 656
<212> PRT
<213> Saccharomyces cerevisiae
<400> 188
Met Lys Val Leu Cys Val Ala Glu Lys Asn Ser Ile Ala Lys Ala Val
Ser Gln Ile Leu Gly Gly Gly Arg Ser Thr Ser Arg Asp Ser Gly Tyr
                                 25
Met Tyr Val Lys Asn Tyr Asp Phe Met Phe Ser Gly Phe Pro Phe Ala
                             40
         35
Arg Asn Gly Ala Asn Cys Glu Val Thr Met Thr Ser Val Ala Gly His
     50
                         55
                                             60
Leu Thr Gly Ile Asp Phe Ser His Asp Ser His Gly Trp Gly Lys Cys
 65
                     70
                                         75
Ala Ile Gln Glu Leu Phe Asp Ala Pro Leu Asn Glu Ile Met Asn Asn
                 85
                                     90
Asn Gln Lys Lys Ile Ala Ser Asn Ile Lys Arg Glu Ala Arg Asn Ala
                                105
            100
```

Asp Tyr Leu Met Ile Trp Thr Asp Cys Asp Arg Glu Gly Glu Tyr Ile

115

120

125

Gly Trp Glu Ile Trp Gln Glu Ala Lys Arg Gly Asn Arg Leu Ile Gln 130 135 140

Asn Asp Gln Val Tyr Arg Ala Val Phe Ser His Leu Glu Arg Gln His 145 150 155 160

Ile Leu Asn Ala Ala Arg Asn Pro Ser Arg Leu Asp Met Lys Ser Val 165 170 175

His Ala Val Gly Thr Arg Ile Glu Ile Asp Leu Arg Ala Gly Val Thr 180 185 190

Phe Thr Arg Leu Leu Thr Glu Thr Leu Arg Asn Lys Leu Arg Asn Gln
195 200 205

Ala Thr Met Thr Lys Asp Gly Ala Lys His Arg Gly Gly Asn Lys Asn 210 220

Asp Ser Gln Val Val Ser Tyr Gly Thr Cys Gln Phe Pro Thr Leu Gly
225 230 235 240

Phe Val Val Asp Arg Phe Glu Arg Ile Arg Asn Phe Val Pro Glu Glu 245 250 255

Phe Trp Tyr Ile Gln Leu Val Val Glu Asn Lys Asp Asn Gly Gly Thr 260 265 270

Thr Thr Phe Gln Trp Asp Arg Gly His Leu Phe Asp Arg Leu Ser Val 275 280 285

Leu Thr Phe Tyr Glu Thr Cys Ile Glu Thr Ala Gly Asn Val Ala Gln 290 295 300

Val Val Asp Leu Lys Ser Lys Pro Thr Thr Lys Tyr Arg Pro Leu Pro 305 310 315 320

Leu Thr Thr Val Glu Leu Gln Lys Asn Cys Ala Arg Tyr Leu Arg Leu 325 330 335

Asn Ala Lys Gln Ser Leu Asp Ala Ala Glu Lys Leu Tyr Gln Lys Gly 340 345 350

Phe Ile Ser Tyr Pro Arg Thr Glu Thr Asp Thr Phe Pro His Ala Met 355 360 365

Asp Leu Lys Ser Leu Val Glu Lys Gln Ala Gln Leu Asp Gln Leu Ala

370 375 380

Ala Gly Gly Arg Thr Ala Trp Ala Ser Tyr Ala Ala Ser Leu Leu Gln 385 390 395 400

Pro Glu Asn Thr Ser Asn Asn Lys Phe Lys Phe Pro Arg Ser Gly 405 410 415

Ser His Asp Asp Lys Ala His Pro Pro Ile His Pro Ile Val Ser Leu 420 425 430

Gly Pro Glu Ala Asn Val Ser Pro Val Glu Arg Arg Val Tyr Glu Tyr 435 440 445

Val Ala Arg His Phe Leu Ala Cys Cys Ser Glu Asp Ala Lys Gly Gln 450 455 460

Ser Met Thr Leu Val Leu Asp Trp Ala Val Glu Arg Phe Ser Ala Ser 465 470 475 480

Gly Leu Val Val Leu Glu Arg Asn Phe Leu Asp Val Tyr Pro Trp Ala 485 490 495

Arg Trp Glu Thr Thr Lys Gln Leu Pro Arg Leu Glu Met Asn Ala Leu 500 505 510

Val Asp Ile Ala Lys Ala Glu Met Lys Ala Gly Thr Thr Ala Pro Pro 515 520 525

Lys Pro Met Thr Glu Ser Glu Leu Ile Leu Leu Met Asp Thr Asn Gly 530 535 540

Ile Gly Thr Asp Ala Thr Ile Ala Glu His Ile Asp Lys Ile Gln Val 545 550 555 560

Arg Asn Tyr Val Arg Ser Glu Lys Val Gly Lys Glu Thr Tyr Leu Gln 565 570 575

Pro Thr Thr Leu Gly Val Ser Leu Val His Gly Phe Glu Ala Ile Gly 580 585 590

Leu Glu Asp Ser Phe Ala Lys Pro Phe Gln Arg Arg Glu Met Glu Gln 595 600 605

Asp Leu Lys Lys Ile Cys Glu Gly His Ala Ser Lys Thr Asp Val Val 610 615 620

Lys Asp Ile Val Glu Lys Tyr Arg Lys Tyr Trp His Lys Thr Asn Ala

625 630 635 640

Cys Lys Asn Thr Leu Leu Gln Val Tyr Asp Arg Val Lys Ala Ser Met 645 650 655

```
<211> 1937
<212> DNA
<213> Saccharomyces cerevisiae
<400> 189
cttcgttcac acttaatatt aaataacagc tctttcctaa taataacata tacactagat 60
atataatacc aaataaatta aaaaaaaaga aaaacaaaaa ataacgtagc tttgttacag 120
tcgtaaaaaa aagaacaata aaccaatctt attgccagcg tctaactagt cctattatat 180
tccaatatat taaggggtaa ggactactat tattcgccct gaattgaaat cttttagaaa 240
gcacctgttc tctctctggt gttcttttt tctcatctat tatctaattt cttcaacctt 300
cgttatttgt tgttattccg taatcgtggt gctcaacttt tgaaatttca cttgtttacc 360
ataacggaac ataattaaaa atttgttctt gaaagtcaca agcactacgt gaacacaaaa 420
ttaaggcaag agtgacaaag taaccctcac aaggagcctg tcgttggtta tattggaagc 480
tatagatata atcgaatcca atgactggtc ctggacctga aataaataag gaggagcacc 540
ccagttctcc gggcaagaaa cagataacat ataatagtat acccaagaat gcaaatctaa 600
ttgatggatc tacgaattca tcgaagaggc caattgaaaa gtatgacaag agaatagccg 660
acccaacaaa aagctatttt ccacatagca tatcaagaac accaaggaga aaatatactt 720
acattctagt cctcacatca ctaaatggaa cttttgagag caaacatgtg gtgataccat 780
ttaaaccaga tggtttgaaa ttgggaaggc ctgttgctaa tagtaatagc agctccagtt 840
catcgctcag gggcggtaaa agagtggatt cacacacttt ttcccaagta aggtccgata 900
atggtaattt cgattcaaga gtactctcca ggaatcatgc attgttaagt tgcgacccgc 960
tcacgggtaa ggtatatata cgagacttaa aatccagtaa tggtacgttc attaacggtc 1020
aaagaattgg ctcaaatgac gtagaaatca aagttggcga tgtgatagac ttgggaacag 1080
atatagatac gaaaattgag catcgaaaaa taagtgccac agttgaagaa ctatttgtac 1140
aacctttatt agaatcaccg atttttgaaa atgaagatag tgatgattgc catacaatta 1200
cagaaaaaga ggaggctgct gccataacaa gtcacatata tggtgattcc aacaacctag 1260
aattggaaga agttatcctg ggctctgata cagaaatact gagtggaatt tttattaata 1320
attgtattgg tacgagcccg actttatcta atatcatcaa aaccctggca atggaaattc 1380
cattttccaa atgcgataat tttaaattac agtcgatgga gaatttctta attaattata 1440
cgacacatct agaatacacg aacaaacttt tagtggaaaa aaatgatcag caactggtaa 1500
agctgcaaaa tggattaaga aggaaactgt cggggaaata cgaaaagatt atcgaacaaa 1560
```

acagaaatca agtaaaacag ttggaaaggg accatatgtt tttcaaaaag tcatttgaag 1620 tgaagaaaag aagaaataat gaaaagcaaa aaagcatgga aagggaaata gaagacttga 1680 aaactaggtt agaagtggaa cgatataaga attcacaaat gatgaagaag aacaaacaga 1740 aagaacaaga actctcaact gcatctaaaa aaaagactac cgaacatgac actagaggcg 1800 tcccgggcat gaatcctaag ggtactgaca aatttagcat caagaacacg ctatgtaatc 1860 atttcacact attaacatt ggaactattt ccatcgggat tatagctatt gtcttcaaga 1920

236

<210> 189

1937

tcctttcccc caactag

<210> 190

<211> 478 <212> PRT

<213> Saccharomyces cerevisiae

<400> 190

Met Thr Gly Pro Gly Pro Glu Ile Asn Lys Glu Glu His Pro Ser Ser 1 5 10 15

Pro Gly Lys Lys Gln Ile Thr Tyr Asn Ser Ile Pro Lys Asn Ala Asn 20 25 30

Leu Ile Asp Gly Ser Thr Asn Ser Ser Lys Arg Pro Ile Glu Lys Tyr 35 40 45

Asp Lys Arg Ile Ala Asp Pro Thr Lys Ser Tyr Phe Pro His Ser Ile 50 55 60

Ser Arg Thr Pro Arg Arg Lys Tyr Thr Tyr Ile Leu Val Leu Thr Ser 65 70 75 80

Leu Asn Gly Thr Phe Glu Ser Lys His Val Val Ile Pro Phe Lys Pro 85 90 95

Asp Gly Leu Lys Leu Gly Arg Pro Val Ala Asn Ser Asn Ser Ser Ser 100 105 110

Ser Ser Ser Leu Arg Gly Gly Lys Arg Val Asp Ser His Thr Phe Ser 115 120 125

Gln Val Arg Ser Asp Asn Gly Asn Phe Asp Ser Arg Val Leu Ser Arg 130 135 140

Asn His Ala Leu Leu Ser Cys Asp Pro Leu Thr Gly Lys Val Tyr Ile 145 150 155 160

Arg Asp Leu Lys Ser Ser Asn Gly Thr Phe Ile Asn Gly Gln Arg Ile 165 170 175

Gly Ser Asn Asp Val Glu Ile Lys Val Gly Asp Val Ile Asp Leu Gly
180 185 190

Thr Asp Ile Asp Thr Lys Ile Glu His Arg Lys Ile Ser Ala Thr Val 195 200 205

												. 4			
Glu	Glu 210	Leu	Phe	Val	Gln	Pro 215	Leu	Leu	Glu	Ser	Pro 220	Ile	Phe	Glu	Asn
Glu 225		Ser	Asp	Asp	Cys 230	His	Thr	Ile	Thr	Glu 235	Lys	Glu	Glu	Ala	Ala 240
Ala	Ile	Thr	Ser	His 245	Ile	Tyr	Gly	Asp	Ser 250	Asn	Asn	Leu	Glu	Leu 255	Glu
Glu	Val	Ile	Leu 260	Gly	Ser	Asp	Thr	Glu 265	Ile	Leu	Ser	Gly	Ile 270	Phe	Ile
Asn	Asn	Cys 275	Ile	Gly	Thr	Ser	Pro 280	Thr	Leu	Ser	Asn	Ile 285	Ile	Lys	Thr
Leu	Ala 290	Met	Glu	Ile	Pro	Phe 295	Ser	Lys	Cys	Asp	Asn 300	Phe	Lys	Leu	G1n
Ser 305	Met	Glu	Asn	Phe	Leu 310	Ile	Asn	Tyr	Thr	Thr 315	His	Leu	Glu	Tyr	Thr 320
Asn	Lys	Leu	Leu	Val 325	Glu	Lys	Asn	Asp	Gln 330	Gln	Leu	Val	Lys	Leu 335	Gln
Asn	Gly	Leu	Arg 340	Arg	Lys	Leu	Ser	Gly 345	Lys	Tyr	Glu	Lys	Ile 350	Ile	Glu
Gln	Asn	Arg 355	Asn	Gln	Val	Lys	Gln 360	Leu	Glu	Arg	Asp	His 365	Met	Phe	Phe
Lys	Lys 370	Ser	Phe	Glu	Val	Lys 375	Lys	Arg	Arg	Asn	Asn 380	Glu	Lys	Gln	Lys
Ser 385	Met	Glu	Arg	Glu	Ile 390	Glu	Asp	Leu	Lys	Thr 395	Arg	Leu	Glu	Val	Glu 400
Arg	Tyr	Lys	Asn	Ser 405	Gln	Met	Met	Lys	Lys 410	Asn	Lys	Gln	Lys	Glu 415	Gln
Glu	Leu	Ser	Thr 420	Ala	Ser	Lys	Lys	Lys 425	Thr	Thr	Glu	His	Asp 430	Thr	Arg
Gly	Val	Pro 435	Gly	Met	Asn	Pro	Lys 440	Gly	Thr	Asp	Lys	Phe 445	Ser	Ile	Lys

238

455

Asn Thr Leu Cys Asn His Phe Thr Leu Leu Thr Phe Gly Thr Ile Ser

460





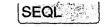
Ile Gly Ile Ile Ala Ile Val Phe Lys Ile Leu Ser Pro Asn 465 470 475

<210> 191 <211> 2849 <212> DNA <213> Saccharomyces cerevisiae

## <400> 191

cgaacatacc tgatatcaga atagatccca atggtgctat attacgtgta agagagagat 60 ttaatttgaa tatgtccgaa gaagatgcca cagtgcattt tcagaatcta atcaatgata 120 gtgtaaatgc tttgctgcct atcgtgattg atcatttaca taatctggca caatactggc 180 ggacctgatt ggttgataat tggtgcttca aaatttaaat ttcgtcactc taattatact 240 taacatataa atggtacctt caatatatct tcttagtaaa aagtagcatg attttgtgtt 300 aatttgcact ttttttattt aaaaataaaa atcacagtta atttttcatg atcttgcaaa 360 gacacgcctc cccctaatgt ggcatatata acaattgtga atcagaaaaa ctcaacactt 420 taacataatg gcgggcacga aggctaaaca aacaagatta gcattgaatg ccttttttgg 480 gtacaataga acagtactga atgacatcat atategaaag geteaagteg geageategt 540 atcttgatac agttccagat gagcatcatg atttcagaaa acccaccgcc aaggttgtaa 600 cgacgcaact gactattgct acttcactag gtatttttgc tttgctttcg ttctcaattc 660 tactaaaaaa gtggcctaga ttatacgcaa gcagacgata taaagatgac gggaaccttc 720 gcttaccgtc ctggaatcag tcaagtttat tcggctggtt aacagtgttg tataagatac 780 gggacgaaca gattctggaa tatgcaggtt tagatgcgta tgtgtttttg agttttttca 840 aaatgtgcat caaattactt tetatttttt gtttettete tgtgtgcgtt atateteeag 900 taagatatca ttttactgga aaaattgatg acggcaacga tgatgatgac agtgaaagtt 960 ctttaataca tctggtaaaa agaattgtgg agggaagtgg cgatggtgac aatcattcag 1020 ctcccgaacg tacaaatgtg tacctctgga tgtatgttct cttcacatat ttcttcacct 1080 tcatagcaat taaaatggca gttgcggaaa caaagcacgt cgtaagtact aggcaagctt 1140 atcttgggaa gcaaaatacg attactgata gaacgataag gctctcaggc atcccgatag 1200 agcttcgtga ttcggaagcc ttaaagacca gaattgaaca attaaaaatc ggcaccgttt 1260 catcaatcac tatttgtcga gagtggggtc ctttgaacaa gctatttcat tgtcggaaga 1320 aaatactcaa aaacctggaa ttaaaatatt cagaatgtcc aagggagctc cgtactcgac 1380 aaccatattc ggagaactat catttattgg gaaacgagca atcaggcgca gttacacatg 1440 gagaaaatgt tccatcaagc aataataatg acgaggatac tatactatat tctcaaattt 1500 ctcttggaga gagaccaaaa atgaaaattg gatatcgtgg tatctttggg aaggaagtag 1560 atgccataga gtacctggag cagcaattaa aatttattga tgctgaaatt attgaagcga 1620 ggaaacaaca ctactccgca acacctacgg cattcgttac gatggattct gttgctaatg 1680 cgcaaatggc agctcaggcg gtattagatc ctagagtaca ttacttcata accagattgg 1740 ctcctgcacc tcacgatatc aagtgggatc atgtttgtct ttctagaaag gaccggttaa 1800 caaaagttta ttctactacc gtatttatag gcctttcaag tttgttttta gtcattcctg 1860 tatcatactt agccacattg ctaaatttga aaacactttc gaaattctgg ccaagcgtag 1920 ggcagctact aaaagatcac cagtgggccg ctaacattgt aacggggcta ttaccaacct 1980 atctcttcac attgcttaac tttggaatcc cctattttta cgagtatttg acttcttatc 2040 aaggattggt atcatacagc gaagaggaaa tctcacttgt ttcaaaaaaat ttcttttata 2100 tttttgtcaa cettttetta gttttcacat tggcaggtac agcatetaat tactgggcgt 2160 acctcagcga taccaccaaa attgcttatc aacttgctac atctgtgaag gagttctcct 2220

Printed:17-08-2000 239



```
tattctatgt cgatttgatt atattgcaag gtattggtat gttcccgftt aagttgttat 2280 tagttggtag tttgatcggc tttcctctag tgaaaatcaa ggctaagaca cctaggcaac 2340 ggaatgaact ttacaatcca ccgatattta actttggact acaattacca cagccaattc 2400 tgattttgat tataacgttg atctacagtg taatgagcac gaaaattttg acttcagggc 2460 tggcgtactt tattattggg ttttacgtct ataaatatca attgatttt gccacagatc 2520 atttgcccca ttctacagga aaagtatggc cattaatttt tagaagaatc atcgttggat 2580 tgctattgt tcaattaca atgacaggaa cactggcagg attcgaagga ggatgggttt 2640 tgtcatcttg cctttcccg cttccagtag tgacattatg tttcctatat gatttcgaga 2700 agaactattt gccgttgtca aaatatatcg cattgagttc aattcgcgag tacgaaagag 2760 acaattctac ggtgaattct gccaatgagg aagagtcgta tgcataccct tacgctgtga 2820 gtgaattaga gggtccgatg ttggattga
```

```
<210> 192
```

<211> 782

<212> PRT

<213> Saccharomyces cerevisiae

<400> 192

Met Thr Ser Tyr Ile Glu Arg Leu Lys Ser Ala Ala Ser Tyr Leu Asp 1 5 10 15

Thr Val Pro Asp Glu His His Asp Phe Arg Lys Pro Thr Ala Lys Val 20 25 30

Val Thr Thr Gln Leu Thr Ile Ala Thr Ser Leu Gly Ile Phe Ala Leu
35 40 45

Leu Ser Phe Ser Ile Leu Leu Lys Lys Trp Pro Arg Leu Tyr Ala Ser 50 55 60

Arg Arg Tyr Lys Asp Asp Gly Asn Leu Arg Leu Pro Ser Trp Asn Gln 65 70 75 80

Ser Ser Leu Phe Gly Trp Leu Thr Val Leu Tyr Lys Ile Arg Asp Glu 85 90 95

Gln Ile Leu Glu Tyr Ala Gly Leu Asp Ala Tyr Val Phe Leu Ser Phe 100 105 110

Phe Lys Met Cys Ile Lys Leu Leu Ser Ile Phe Cys Phe Phe Ser Val 115 120 125

Cys Val Ile Ser Pro Val Arg Tyr His Phe Thr Gly Lys Ile Asp Asp 130 135 140

Gly Asn Asp Asp Asp Ser Glu Ser Ser Leu Ile His Leu Val Lys 145 150 155 160



- Arg Ile Val Glu Gly Ser Gly Asp Gly Asp Asn His Ser Ala Pro Glu 165 170 175
- Arg Thr Asn Val Tyr Leu Trp Met Tyr Val Leu Phe Thr Tyr Phe Phe 180 185 190
- Thr Phe Ile Ala Ile Lys Met Ala Val Ala Glu Thr Lys His Val Val
  195 200 205
- Ser Thr Arg Gln Ala Tyr Leu Gly Lys Gln Asn Thr Ile Thr Asp Arg 210 215 220
- Thr Ile Arg Leu Ser Gly Ile Pro Ile Glu Leu Arg Asp Ser Glu Ala 225 230 235 240
- Leu Lys Thr Arg Ile Glu Gln Leu Lys Ile Gly Thr Val Ser Ser Ile 245 250 255
- Thr Ile Cys Arg Glu Trp Gly Pro Leu Asn Lys Leu Phe His Cys Arg 260 265 270
- Lys Lys Ile Leu Lys Asn Leu Glu Leu Lys Tyr Ser Glu Cys Pro Arg 275 280 285
- Glu Leu Arg Thr Arg Gln Pro Tyr Ser Glu Asn Tyr His Leu Leu Gly 290 295 300
- Asn Glu Gln Ser Gly Ala Val Thr His Gly Glu Asn Val Pro Ser Ser 305 310 315 320
- Asn Asn Asn Asp Glu Asp Thr Ile Leu Tyr Ser Gln Ile Ser Leu Gly 325 330 335
- Glu Arg Pro Lys Met Lys Ile Gly Tyr Arg Gly Ile Phe Gly Lys Glu
- Val Asp Ala Ile Glu Tyr Leu Glu Gln Gln Leu Lys Phe Ile Asp Ala 355 360 365
- Glu Ile Ile Glu Ala Arg Lys Gln His Tyr Ser Ala Thr Pro Thr Ala 370 375 380
- Phe Val Thr Met Asp Ser Val Ala Asn Ala Gln Met Ala Ala Gln Ala 385 390 395 400
- Val Leu Asp Pro Arg Val His Tyr Phe Ile Thr Arg Leu Ala Pro Ala 405 410 415



Pro His Asp Ile Lys Trp Asp His Val Cys Leu Ser Arg Lys Asp Arg
420 425 430

Leu Thr Lys Val Tyr Ser Thr Thr Val Phe Ile Gly Leu Ser Ser Leu 435 440 445

Phe Leu Val Ile Pro Val Ser Tyr Leu Ala Thr Leu Leu Asn Leu Lys 450 455 460

Thr Leu Ser Lys Phe Trp Pro Ser Val Gly Gln Leu Leu Lys Asp His 465 470 475 480

Gln Trp Ala Ala Asn Ile Val Thr Gly Leu Leu Pro Thr Tyr Leu Phe
485 490 495

Thr Leu Leu Asn Phe Gly Ile Pro Tyr Phe Tyr Glu Tyr Leu Thr Ser 500 505 510

Tyr Gln Gly Leu Val Ser Tyr Ser Glu Glu Glu Ile Ser Leu Val Ser 515 520 525

Lys Asn Phe Phe Tyr Ile Phe Val Asn Leu Phe Leu Val Phe Thr Leu 530 535 540

Ala Gly Thr Ala Ser Asn Tyr Trp Ala Tyr Leu Ser Asp Thr Thr Lys 545 550 555 560

Ile Ala Tyr Gln Leu Ala Thr Ser Val Lys Glu Phe Ser Leu Phe Tyr 565 570 575

Val Asp Leu Ile Ile Leu Gln Gly Ile Gly Met Phe Pro Phe Lys Leu 580 585 590

Leu Leu Val Gly Ser Leu Ile Gly Phe Pro Leu Val Lys Ile Lys Ala 595 600 605

Lys Thr Pro Arg Gln Arg Asn Glu Leu Tyr Asn Pro Pro Ile Phe Asn 610 615 620

Phe Gly Leu Gln Leu Pro Gln Pro Ile Leu Ile Leu Ile Ile Thr Leu 625 630 635 640

Ile Tyr Ser Val Met Ser Thr Lys Ile Leu Thr Ser Gly Leu Ala Tyr
645 650 655

Phe Ile Ile Gly Phe Tyr Val Tyr Lys Tyr Gln Leu Ile Phe Ala Thr 660 665 670

```
Asp His Leu Pro His Ser Thr Gly Lys Val Trp Pro Leu Ile Phe Arg
        675
                            680
Arg Ile Ile Val Gly Leu Leu Phe Gln Leu Thr Met Thr Gly Thr
                        695
                                            700
Leu Ala Gly Phe Glu Gly Gly Trp Val Leu Ser Ser Cys Leu Phe Pro
                    710
                                        715
                                                             720
Leu Pro Val Val Thr Leu Cys Phe Leu Tyr Asp Phe Glu Lys Asn Tyr
                                    730
                725
                                                         735
Leu Pro Leu Ser Lys Tyr Ile Ala Leu Ser Ser Ile Arg Glu Tyr Glu
            740
                                745
                                                     750
Arg Asp Asn Ser Thr Val Asn Ser Ala Asn Glu Glu Glu Ser Tyr Ala
        755
                            760
                                                765
```

Tyr Pro Tyr Ala Val Ser Glu Leu Glu Gly Pro Met Leu Asp

780

775

<210> 193

770

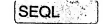
<211> 1781

<212> DNA

<213> Saccharomyces cerevisiae

<400> 193

ttcaagtgca ctctttgtaa ggaagattac aagtctccag tggtcacaaa ctgtggccat 60 tatttttgtg gatcgtgttt tgccaaagat atgaaaaagg gcaccaaatg ctttatatgc 120 cacaaggaga cccacggcag tgcaaaagta gcatctgact tacagaagat gctaaataaa 180 aggaaaagtt gatggagaat gccatatcac cccaaaaatt acacgcaccc gatgctaatg 240 tacaggaatt atagagcaca tgacccatag atttatcgag cattgttgca atttcgaaag 300 actettteae ataataaagt atgtaaacta tatagataga agatgteeeg tgtetttttg 360 tetactaaat gatgatetge teatttaaag tegeegegae taetttgaca aaaaaaaaaa 420 acttagaaaa tacgacaaat agagattatt gaatgaagta cattgaaaaa agaaagaaga 480 aaggcacata gcagcacaca atgtcgcacc aaaaccagct tattccacaa gcttatattt 540 ctaactttca taacagattg acaaacgaag atgatggtat ccccatcttt acaatggctc 600 aacaaacaag gcagcataaa agggctaaag tggtcaacta tgcggaatat gacaacgatc 660 tetttgatga atteaatatg aaeggtteta attteaacaa tgetgataca caetataaag 720 ataatgcagt gtctcatgaa aatactccgg cacttacaaa tggtgttacc atggacggtt 780 ccgaatacaa tgtcctagag aacatgaatg gagctgatag tattatctct aacaacaaat 840 acgatgcggg ttcaaacatg gttgtggaat ctttatccgg tttgaatagc aataacaacg 900 ccagcaatgg tccgagcaac aaagcgcagg cacaggatat tggaaacgcc gttctaccgg 960 atctgcaaga ccaacaccac aaccccttca acatattgag ataccctaaa ataagagata 1020 ctttcattaa cggaaaagtg gtgtctccat atagactcaa cactgatcaa gaaacgaagg 1080



```
caaacgccaa ttctggagag gcaatcatga taccaattac ttttggatata gaacatatgg 1140 gtcataccat aaaagaccag tttctctgga actacaatga cgactccata tctccggagg 1200 aatttgcctc tatatactgt aaagatcttg atatgacttc cgctacctta caaactcaaa 1260 ttgcgaatat aataaaagag cagttgaaag acctcgaaaa tattgcagcc actgagataa 1320 tgtctgacct ccacgtgata atcaacctaa cctgcaactt acaagacaga ttttttgaag 1380 ataacttcca gtggaacttg aacgacaaat cacttactcc agaaagattt gctacatcca 1440 ttgtacagga ccttggctta acaagagagt tcatcccctt aatatctcaa tcgcttcatg 1500 aaactatctt gaagataaag aaggactggg tagatggcca cttgattcag gaccatgtcc 1560 caaacgatgc cgcatttggg tacttatctg gtataaggct ggatattgat gaactgggct 1620 ccaattggtg cccaagggtg gaaatattaa caaaagaaga aaacttaaga agattgaaaa gagaaactga tagattatct agaaggggca 1740 ggagaagatt agatgacta gaaccacaa tgagaatgta g
```

```
<210> 194
```

<211> 426

<212> PRT

<213> Saccharomyces cerevisiae

<400> 194

Met Ser His Gln Asn Gln Leu Ile Pro Gln Ala Tyr Ile Ser Asn Phe

1 5 10 15

His Asn Arg Leu Thr Asn Glu Asp Asp Gly Ile Pro Ile Phe Thr Met 20 25 30

Ala Gln Gln Thr Arg Gln His Lys Arg Ala Lys Val Val Asn Tyr Ala 35 40 45

Glu Tyr Asp Asn Asp Leu Phe Asp Glu Phe Asn Met Asn Gly Ser Asn 50 55 60

Phe Asn Asn Ala Asp Thr His Tyr Lys Asp Asn Ala Val Ser His Glu
65 70 75 80

Asn Thr Pro Ala Leu Thr Asn Gly Val Thr Met Asp Gly Ser Glu Tyr 85 90 95

Asn Val Leu Glu Asn Met Asn Gly Ala Asp Ser Ile Ile Ser Asn Asn 100 105 110

Lys Tyr Asp Ala Gly Ser Asn Met Val Val Glu Ser Leu Ser Gly Leu 115 120 125

Asn Ser Asn Asn Asn Ala Ser Asn Gly Pro Ser Asn Lys Ala Gln Ala 130 135 140

Gln Asp Ile Gly Asn Ala Val Leu Pro Asp Leu Gln Asp Gln His His

													,		
145					150					155					160
Asn	Pro	Phe	Asn	Ile 165	Leu	Arg	Tyr	Pro	Lys 170	Ile	Arg	Asp	Thr	Phe 175	Ile
Asn	Gly	Lys	Val 180	Val	Ser	Pro	Tyr	Arg 185	Leu	Asn	Thr	Asp	Gln 190	Glu	Thr
Lys	Ala	Asn 195	Ala	Asn	Ser	Gly	Glu 200	Ala	Ile	Met	Ile	Pro 205	Ile	Thr	Leu
Asp	Ile 210	Glu	His	Met	Gly	His 215	Thr	Ile	Lys	Asp	Gln 220	Phe	Leu	Trp	Asn
Tyr 225	Asn	Asp	Asp	Ser	Ile 230	Ser	Pro	Glu	Glu	Phe 235	Ala	Ser	Ile	Tyr	Cys 240
Lys	Asp	Leu	Asp	Met 245	Thr	Ser	Ala	Thr	Leu 250	Gln	Thr	Gln	Ile	Ala 255	Asn
Ile	Ile	Lys	Glu 260		Leu	Lys	Asp	Leu 265	Glu	Asn	Ile	Ala	Ala 270	Thr	Glu
Ile	Met	Ser 275		Leu	His	Val	Ile 280		Asn	Leu	Thr	Cys 285	Asn	Leu	Gln
Asp	Arg 290		Phe	e Glu	Asp	Asn 295	Phe	Gln	Trp	) Asn	Leu 300	Asn	Asp	Lys	Ser
Lev		Pro	Glu	a Arg	Phe		Thr	Ser	: Ile	e Val	Gln S	Asp	Let	ı Gly	7 Leu 320
Thi	. Arg	g Glu	ı Phe	325		Lev	ılle	e Ser	Glr 330	n Sei	r Leu	His	s Glı	335	c Ile
Le	ı Lys	s Ile	E Lys		s Ası	Tr	val	L Asp		y Hi:	s Lev	ı Ile	35	n Ası O	p His
Va:	l Pro	35!		o Ala	a Ala	a Phe	360	у Туз )	r Le	u Se	r Gly	y Ile 36	e Ar	g Le	u Asp
11	e Asj		u Lei	u Gl	y Se	r Ası		р Су:	s Pr	o Ar	g Va 38	1 Gl <sup>.</sup> 0	u Il	e Le	u Thr

245

Lys Glu Glu Ile Gln Lys Arg Glu Ile Glu Lys Glu Arg Asn Leu Arg

Arg Leu Lys Arg Glu Thr Asp Arg Leu Ser Arg Arg Gly Arg Arg Arg

390

395

370

405

410

415

Leu Asp Asp Leu Glu Thr Thr Met Arg Met 420 425

<210> 195

<211> 815

<212> DNA

<213> Saccharomyces cerevisiae

<400> 195

gtcatatcaa gatcttaca gtatatagag gcaaattcct ccggagatat ggagtcgtca 60 ttgtagttcc agagaaactg gtctttatg gtatgaccca tatgttctat atccaaagta 120 attggtatca tgattgcctc tccagaattg gcgtttgcct tcgtttcttg atcagtgttg 180 agtctatatg gagacaccac ttttccgtta atgaaagtat ctcttatttt agggtatctc 240 aatatgttga aggggttgtg gtgttggtct tgcagatccg gtagaacggc gtttccaata 300 tcctgtgcct gcgctttgtt gctcggacca ttgctggcgt tgttattgct attcaaaccg 360 gataaagatt ccacaaccat gtttgaaccc gcatcgtatt tgttgttaga gataatacta 420 tcagctccat tcatgttctc taggacattg tattcggaac cgtccatggt aacaccattt 480 gtaagtgccg gagtatttc atgaatca tcaaagagat cgttgtcata ttccgcatag 600 ttgaccactt tagcccttt atgactcat attgattga ccattgtaaa gatggggata 660 ccatcatctt tagcccttt atgttataga aagttagaa tataagcttg tggaataagc 720 tggttttggt gcgacattgt gtgctgtat gtgccttct tcttcttt ttcaatgtac 780 ttcattcaat aatcctatt tgtcgtatt tctaa

<210> 196

<211> 104

<212> PRT

<213> Saccharomyces cerevisiae

<400> 196

Met Arg His Cys Ile Ile Phe Ile Val Cys Ile Ser Ile Val Glu Ile 1 5 10 15

Arg Thr Val His Ile Glu Phe Ile Lys Glu Ile Val Val Ile Phe Arg
20 25 30

Ile Val Asp His Phe Ser Pro Phe Met Leu Pro Cys Leu Leu Ser His
35 40 45

Cys Lys Asp Gly Asp Thr Ile Ile Phe Val Cys Gln Ser Val Met Lys 50 55 60

Val Arg Asn Ile Ser Leu Trp Asn Lys Leu Val Leu Val Arg His Cys

```
65 70 75 80
```

Val Leu Cys Ala Phe Leu Leu Ser Phe Phe Asn Val Leu His Ser 85 90 95

Ile Ile Ser Ile Cys Arg Ile Phe 100

```
<210> 197
<211> 737
```

<212> DNA

<213> Saccharomyces cerevisiae

```
<400> 197
```

ccacatytca caactacttt gtgaagttgc aatgegtgat tagtattata aaacatcata 60 geettgecaa ataaactege tteecagaaa agaegeeaaa ttaaetgeeg etgttatytg 120 acaaaacagg geateteaca tattegegta etggtgteet tttageteat teegatatta 180 tteeaagaag gaattteeat etagagaaaa tgeateegtg catttteata aaceeacaca 240 attaaaatge ettgegaaaa ggaggaeteg teegtgeaac tgttgaaaaa aatagaegga 300 geateatacg ttegagtgga aaattatgga gagttteea agetetatgg catgtagagt 360 egtgattget getgtaeget tttgeacaat attgaateet eaatetaaag aattaaatte 420 tetaatteea atgtagaaat attteaaetg ttagttttt attteaggtt gaatatagta 480 egacaaaata teaaggaaaa atggetagag aaateacega cateaaacaa tttttggaat 540 tggetaagee atteagaea aceaagttea aggttagagg etettetet ttgtaeact 660 tggttateaa egatgetggt aaggetaaga aattgateea atetttgeea ceaactttga 720 aggttaacag attaaa

```
<210> 198
```

<211> 78

<212> PRT

<213> Saccharomyces cerevisiae

## <400> 198

Met Ala Arg Glu Ile Thr Asp Ile Lys Gln Phe Leu Glu Leu Thr Arg

1 5 10 15

Arg Ala Asp Val Lys Thr Ala Thr Val Lys Ile Asn Lys Lys Leu Asn 20 25 30

Lys Ala Gly Lys Pro Phe Arg Gln Thr Lys Phe Lys Val Arg Gly Ser 35 40 45

Ser Ser Leu Tyr Thr Leu Val Ile Asn Asp Ala Gly Lys Ala Lys Lys 50 55 60

Leu Ile Gln Ser Leu Pro Pro Thr Leu Lys Val Asn Arg Leu

```
65
                      70
                                          75
<210> 199
<211> 1331
<212> DNA
<213> Saccharomyces cerevisiae
<400> 199
taagggctgt caatttttca tttccctcaa agttaatgcg caagagttag gatgatcttg 60
attattagtt atttgtttgt tccatcaatt atgccgacgt ccatgaatgc tattgaagta 120
tcgattactt atgctattct attgtatttg agccggcggc tagtaaacaa gacagcatac 180
ctttccaaca tccgtacata ttgattattg tatctatgat tttcatgtaa tgtatgggtt 240
aactaatcct ttctcttctt ttttttttt ctgaaaatca gtagctagag gaacctgttc 300
gtgaaagata tggaatattc cgtctaggga agagccgaga gctgagagca gtatccatcg 360
gatggaatct tgcctggatg cgtaacgacg acagcttgtc tagaaggcga agtacaagat 420
actaagaaga ctataacaac agaagacaat tttgtagtgc gattctatat tattgacgta 480
accaacaaaa aagtatcaga atggctaaac aatcattagg tatgtgcaca aatgctaaat 540
cactgaaacc ctgtagagct ggaattgaaa atatagtttg taccttttgc ctgagggcag 600
aatgactcaa tgttatctta attcgattag aaaaagaagg tgcgttatcc aaggttgaaa 660
cttgtaaact atttaggtga tgtctcataa ctgcaaccac tggaaaatga cgcaaatatc 720
aaaataatag ggagataaag totcacgaca tgaaaaaatc atagaattat tgttcaattt 780
atctagccgt atcttacctt atggaaaaag tgtgaatatg tttttctaaa tcgagcttcc 840
cggaaattac actatattat gcatccgtga tacatctgaa attcaatact ttctccaagg 900
tcctgcttac taagagggta agatttgttg aaactcgaaa tactaacatt attttcatta 960
ttttagacgt ttcctccgac agaagaaagg ctagaaaggc ttatttcacc gccccatcct 1020
cccaacgtcg tgttttgcta tctgctccat tatctaagga gttgagagct caatacggta 1080
tcaaggcctt gccaatcaga agagatgatg aagttttggt tgttcgtggt tccaagaagg 1140
gtcaagaagg taagatttca tetgtttaca gattgaaatt tgccgttcaa gttgacaagg 1200
ttaccaagga aaaggtcaac ggtgcttccg ttccaattaa cttgcaccca tccaagcttg 1260
tcattactaa attacatttg gacaaggaca gaaaagcttt gatccaaaga aagggcggta 1320
agttggaata a
                                                                   1331
<210> 200
<211> 127
<212> PRT
<213> Saccharomyces cerevisiae
<400> 200
Met Ala Lys Gln Ser Leu Asp Val Ser Ser Asp Arg Arg Lys Ala Arg
                                     10
Lys Ala Tyr Phe Thr Ala Pro Ser Ser Gln Arg Arg Val Leu Leu Ser
                                 25
```

```
Ala Pro Leu Ser Lys Glu Leu Arg Ala Gln Tyr Gly Ile Lys Ala Leu
                                                  45
         35
                              40
Pro Ile Arg Arg Asp Asp Glu Val Leu Val Val Arg Gly Ser Lys Lys
                                              60
                         55
     50
Gly Gln Glu Gly Lys Ile Ser Ser Val Tyr Arg Leu Lys Phe Ala Val
                     70
 65
Gln Val Asp Lys Val Thr Lys Glu Lys Val Asn Gly Ala Ser Val Pro
                                      90
                 85
Ile Asn Leu His Pro Ser Lys Leu Val Ile Thr Lys Leu His Leu Asp
                                 105
            100
Lys Asp Arg Lys Ala Leu Ile Gln Arg Lys Gly Gly Lys Leu Glu
                             120
        115
<210> 201
```

```
<211> 1376
<212> DNA
<213> Saccharomyces cerevisiae
```

<400> 201 actttttgtc tgctgctggt cgtttgtctt tcgttttaaa attgcgctag acaagtaaac 60 agggattgct taagaatcaa agtagcttaa ctctaaagta ttattttcct cagttgtggg 120 cccatgtgtt ggagggaagg aatatattga aatgtaaatg ttcttaagtt cggttgaact 180

tggatattgt tacaagagtt ctagtctttg ataccatttt tacgcaatta caaccgcatt 240 atttaccttt tcatcttcag ttttacggtt cagtttattc tgttacgaaa gaactatggt 300 gattcaaagg cgaagtgcgt aggattgtaa ctcctatatc tttaggatac ttacaatttt 360 gtactgtttt caagaccact gtaaccgata ataaaccgga ggacacattt taacccacta 420 tttttttcag aagatcagat gcgagagctc gaagcataag tataatacta acgtttcaaa 480 acatagtaat taggtaaaaa atgactcgct cttccgtttt agctgatgct ttgaatgcca 540 ttaataacgc cgaaaagacc ggtaaacgtc aggttctatt gagaccttct tccaaggtta 600 tcatcaagtt tttacaagtt atgcaaaagc atggtatgtt ccaactattt ttcaatattt 660 tcacatgtgt ttcaatttct gcttattttt aaatgttacc acgaggtttg tccaagttca 720 atgttgcgca actctaacga agaaataatt attgccattg ttttttactc cgggctgata 780 actagatggt gtgatcgggc agtatactaa tttatactgg acaaagactc gtaaaagatg 840 ttctttgtgc ttagtcccat actgtttttt aagtgtccgg gatatttaat cccatgtgga 900 aatgcttctt acacggttat ggattacacc tcatgtgtag ctactatatc cattaccgtt 960

tacttttcct caaaatctca ctcttaaaat tttcaatggc aaaattcttc cgcacaactt 1020 agacaacatt ttcttgtttt ttatgaagta agcaaaaatt tcgaatcaac aacgctccat 1080 gagattette aatactaaca tttacteett atttaggtta cattggegaa ttegaataca 1140

ttgacgacca cagatctggt aagattgtcg tccaactgaa cggtagattg aacaaatgtg 1200 gtgttatttc cccaagattt aacgttaaga ttggtgacat tgaaaaatgg accgccaact 1260

tattgccagc cagacaattc ggttacgtta ttttgactac ttctgcgggt attatggacc 1320 atgaggaagc tagaagaaag cacgtttccg gtaaaatctt gggttttgtc tattaa 1376

<210> 202

<211> 130

<212> PRT

<213> Saccharomyces cerevisiae

<400> 202

Met Thr Arg Ser Ser Val Leu Ala Asp Ala Leu Asn Ala Ile Asn Asn 1 5 10 15

Ala Glu Lys Thr Gly Lys Arg Gln Val Leu Leu Arg Pro Ser Ser Lys
20 25 30

Val Ile Ile Lys Phe Leu Gln Val Met Gln Lys His Gly Tyr Ile Gly
35 40 45

Glu Phe Glu Tyr Ile Asp Asp His Arg Ser Gly Lys Ile Val Val Gln 50 55 60

Leu Asn Gly Arg Leu Asn Lys Cys Gly Val Ile Ser Pro Arg Phe Asn 65 70 75 80

Val Lys Ile Gly Asp Ile Glu Lys Trp Thr Ala Asn Leu Leu Pro Ala 85 90 95

Arg Gln Phe Gly Tyr Val Ile Leu Thr Thr Ser Ala Gly Ile Met Asp 100 105 110

His Glu Glu Ala Arg Arg Lys His Val Ser Gly Lys Ile Leu Gly Phe 115 120 125

Val Tyr 130

<210> 203

<211> 1340

<212> DNA

<213> Saccharomyces cerevisiae

<400> 203

ccctcaaccc gcattttgct gagaattttc accaaggccc taggtgatat tagattccac 60 ctgactaatt gcattacagc cgacccaagg caatatcagt ttaataaaat atcatgtatc 120 tcaccctctt cttggtatta gtaaagagac gcctgatctt gtaacagtgg tgaagattgt 180

```
actagagcag aatcaagaat ttaaaaagtg taaaggcaag gcagaggcga tgtacataaa 240
acttcgaagt aagaaatatt taatagttct cgccacatca ctatgcagct atataaaaac 300
tactataaac gtttgttttg ttccttacgc acaatatcct tgcctagaaa tcgtttttga 360
aatttaaatt tttattacca tttatttgat tcgccttcag aaaaatatgg aagagtgcat 420
atttaaaaag gactatttca gcatatagta aaagtcaggt tatttgttta tttgcgatat 480
cagagtaact taaactaact atgcagggca cttttaaaaag gttttaccat cccacgctta 540
cgcggatgtc cttcttggat aaattcctca agcctatgat ggcaacggct tccccaaagg 600
aataccagat caaacaactg gtcaagccaa taggcttaac acaagcacca aggaaaagca 660
ccaaatactc ccaggggaac tctttgaggg atatgtttga ttcggaaaag acaaaccaca 720
gagttaaaga gttggccgtt gaattcagca aatctggact ttatgacgtg caagtcttcc 780
aaaagacaaa ggggaaattg tttatagctc cagtttcata ttggaaagaa gataaagctt 840
tgttttttcc tcatttgata ggaacggcaa tggatggtac gaaacaacag aatatcgagg 900
atatgttaag gggtaaaacc agtatagtga ggttatttag tacagcatct ggcgataagt 960
tgagtagttc atacttccaa ggaatcgtag acgataacaa aaaaactgac tacttgactg 1020
aagctgatgc gcgtttaagt ttaaatgaca gtaacgtcca aatcatcgag gtcaatcttg 1080
tagaaaacgc tgtgaaaagt gctctagtga aaacgcttgc tcgttgggcc aatcgcgttc 1140
catcctggcg ccagccattt tatttcgaat gttctagagg ccaatggcca ttttccgtca 1200
gggaagaget ettttgcaat aatgtetttt etggataegt etttettgtg gaceageagt 1260
taaaaattag gtgggcagct tgcggggagg ctactccatc tgaaaaggaa gcattgtgga 1320
                                                                  1340
agtttgccaa acgtctgtga
```

<210> 204

<211> 279

<212> PRT

<213> Saccharomyces cerevisiae

<400> 204

Met Gln Gly Thr Phe Lys Arg Phe Tyr His Pro Thr Leu Thr Arg Met

1 5 10 15

Ser Phe Leu Asp Lys Phe Leu Lys Pro Met Met Ala Thr Ala Ser Pro 20 25 30

Lys Glu Tyr Gln Ile Lys Gln Leu Val Lys Pro Ile Gly Leu Thr Gln 35 40 45

Ala Pro Arg Lys Ser Thr Lys Tyr Ser Gln Gly Asn Ser Leu Arg Asp
50 55 60

Met Phe Asp Ser Glu Lys Thr Asn His Arg Val Lys Glu Leu Ala Val 65 70 75 80

Glu Phe Ser Lys Ser Gly Leu Tyr Asp Val Gln Val Phe Gln Lys Thr 85 90 95

Lys Gly Lys Leu Phe Ile Ala Pro Val Ser Tyr Trp Lys Glu Asp Lys 100 105 110

```
Ala Leu Phe Phe Pro His Leu Ile Gly Thr Ala Met Asp Gly Thr Lys
115 120 125
```

Gln Gln Asn Ile Glu Asp Met Leu Arg Gly Lys Thr Ser Ile Val Arg 130 135 140

Leu Phe Ser Thr Ala Ser Gly Asp Lys Leu Ser Ser Ser Tyr Phe Gln 145 150 155 160

Gly Ile Val Asp Asp Asn Lys Lys Thr Asp Tyr Leu Thr Glu Ala Asp 165 170 175

Ala Arg Leu Ser Leu Asn Asp Ser Asn Val Gln Ile Ile Glu Val Asn 180 185 190

Leu Val Glu Asn Ala Val Lys Ser Ala Leu Val Lys Thr Leu Ala Arg 195 200 205

Trp Ala Asn Arg Val Pro Ser Trp Arg Gln Pro Phe Tyr Phe Glu Cys 210 215 220

Ser Arg Gly Gln Trp Pro Phe Ser Val Arg Glu Glu Leu Phe Cys Asn 225 230 235 240

Asn Val Phe Ser Gly Tyr Val Phe Leu Val Asp Gln Gln Leu Lys Ile 245 250 255

Arg Trp Ala Ala Cys Gly Glu Ala Thr Pro Ser Glu Lys Glu Ala Leu 260 265 270

Trp Lys Phe Ala Lys Arg Leu 275

<210> 205

<211> 1754

<212> DNA

<213> Saccharomyces cerevisiae

<400> 205

ttctattggt gtagttatac cagaaccatc cttcccaagg ccttttcccg caacatatcc 60 catactcgaa agtaacttcg caccaatacc atatgtcttc gttaattttg agatcgttgg 120 ggcattgccg ctactagttt tcaaggcatc attagtgtat gtcatgtcac tattcatgct 180 tgaattatta tcttcctcat cgctataatt ataggaatcg attcttcttt ttttgaaaaa 240 gaacttttta tctgtgttgg agtccgaatc ctccatgttt tggcgtactg aaggcttaag 300 ttcttcgacc tctctggtt gtactattc gaaattttgg atttttgtta ttgttttgac 360

```
ataatgtaaa tactagatgc gcgctctaag gcctcagtat taaaaattgc aagatatccc 420
taacttgata attatttgaa agtcgcataa cgtacgataa attcgatact gcgaggatat 480
tatcaacgta tttaacacct atgaacgaag cagatgttac aaaatttgtt aataatgcca 540
ggaaaacgct gaccgatgct caacttttat gttcaagtgc taatttaagg attgtagata 600
ttaagaaaaa attgtcatct tggcagttga gtatttcaaa actcaatttt ctaatagttg 660
gcttgagaca gcaaggaaag tttctttaca ctattttaaa ggaaggcatt gggacaaagc 720
taatccagaa acaatggaat caagctgtct tagtggtctt agttgatgag atgaagtact 780
ggcaatatga aatcacttct aaagtacaaa ggttggatgg tatagtaaat gaattgagca 840
tatcagaaaa agatgatacc gatccctcta aactaggaga ttacatctcg agagacaacg 900
ttaacttatt gaatgacaaa ctgaaagaag tgccagtaat tgagcgtcaa attgaaaaca 960
ttaagettea ataegaaaat atggteagaa aagttaacaa agaactgatt gataccaagt 1020
tgacggacgt aactcaaaaa ttccaaagta aatttggtat agataacctg atggaaacaa 1080
atgttgcaga gcagtttagc agggaactaa cggaccttga aaaagattta gcagagataa 1140
tgaattcatt gacacagcac tttgataaga cattgctgtt acaagataaa aaaattgata 1200
acgatgaacg tgaggagctg tttaaggtgg tacaaggcga cgacaaagaa ctatacaaca 1260
ttttcaaaac tctgcatgag gtaattgatg acgtggacaa aacaattctt aacttgggtc 1320
aatttttgca ggcaaaaata aaggaaaaga cagaactaca cagcgaagtt tctgaaataa 1380
taaacgattt caatagaaat ttggaatatc tattaatctt taaagatatt tccaatctga 1440
ttgatagett caaaaattee tgtacacaag atattcaaac aactaaggaa etttgtgaat 1500
tttatgataa ttttgaagaa agctacggta acttagttct agaagcaaag aggagaaagg 1560
atgtggcaaa cagaatgaaa actatattga aagattgtga aaagcagttg cagaatttag 1620
atgctcagga ccaggaagaa cgtcagaatt ttatagcgga aaatggaact tatcttcctg 1680
agacaatctg gcccggtaaa attgacgatt tttcttccct gtacacttta aattacaacg 1740
                                                                  1754
tgaagaatcc ttag
```

<210> 206

<211> 417

<212> PRT

<213> Saccharomyces cerevisiae

<400> 206

Met Asn Glu Ala Asp Val Thr Lys Phe Val Asn Asn Ala Arg Lys Thr 1 5 10 15

Leu Thr Asp Ala Gln Leu Leu Cys Ser Ser Ala Asn Leu Arg Ile Val 20 25 30

Asp Ile Lys Lys Leu Ser Ser Trp Gln Leu Ser Ile Ser Lys Leu 35 40 45

Asn Phe Leu Ile Val Gly Leu Arg Gln Gln Gly Lys Phe Leu Tyr Thr 50 55 60

Ile Leu Lys Glu Gly Ile Gly Thr Lys Leu Ile Gln Lys Gln Trp Asn
65 70 75 80

Gln Ala Val Leu Val Val Leu Val Asp Glu Met Lys Tyr Trp Gln Tyr

90

85

95

Glu Ile Thr Ser Lys Val Gln Arg Leu Asp Gly Ile Val Asn Glu Leu 100 105 110

Ser Ile Ser Glu Lys Asp Asp Thr Asp Pro Ser Lys Leu Gly Asp Tyr 115 120 125

Ile Ser Arg Asp Asn Val Asn Leu Leu Asn Asp Lys Leu Lys Glu Val
130 135 140

Pro Val Ile Glu Arg Gln Ile Glu Asn Ile Lys Leu Gln Tyr Glu Asn 145 150 155 160

Met Val Arg Lys Val Asn Lys Glu Leu Ile Asp Thr Lys Leu Thr Asp 165 170 175

Val Thr Gln Lys Phe Gln Ser Lys Phe Gly Ile Asp Asn Leu Met Glu 180 185 190

Thr Asn Val Ala Glu Gln Phe Ser Arg Glu Leu Thr Asp Leu Glu Lys
195 200 205

Asp Leu Ala Glu Ile Met Asn Ser Leu Thr Gln His Phe Asp Lys Thr 210 215 220

Leu Leu Gln Asp Lys Lys Ile Asp Asn Asp Glu Arg Glu Glu Leu 225 230 235 240

Phe Lys Val Val Gln Gly Asp Asp Lys Glu Leu Tyr Asn Ile Phe Lys 245 250 255

Thr Leu His Glu Val Ile Asp Asp Val Asp Lys Thr Ile Leu Asn Leu 260 265 270

Gly Gln Phe Leu Gln Ala Lys Ile Lys Glu Lys Thr Glu Leu His Ser 275 280 285

Glu Val Ser Glu Ile Ile Asn Asp Phe Asn Arg Asn Leu Glu Tyr Leu 290 295 300

Leu Ile Phe Lys Asp Ile Ser Asn Leu Ile Asp Ser Phe Lys Asn Ser 305 310 315 320

Cys Thr Gln Asp Ile Gln Thr Thr Lys Glu Leu Cys Glu Phe Tyr Asp 325 330 335

Asn Phe Glu Glu Ser Tyr Gly Asn Leu Val Leu Glu Ala Lys Arg Arg

340 345 350

Lys Asp Val Ala Asn Arg Met Lys Thr Ile Leu Lys Asp Cys Glu Lys 355 360 365

Gln Leu Gln Asn Leu Asp Ala Gln Asp Gln Glu Glu Arg Gln Asn Phe 370 375 380

Ile Ala Glu Asn Gly Thr Tyr Leu Pro Glu Thr Ile Trp Pro Gly Lys
385 390 395 400

Ile Asp Asp Phe Ser Ser Leu Tyr Thr Leu Asn Tyr Asn Val Lys Asn 405 410 415

Pro

<210> 207

<211> 1342

<212> DNA

<213> Saccharomyces cerevisiae

<400> 207

atctgtaaat aataagccca tgaaatatca atctcacaat gaaagaattg tataattcag 60 aatgatttga cgtaaaagca agtctcattc aagaaactgt aaaatccgta cacactactg 120 aattttacat ccatacattt ttttgaaatt tcatgttttt ttgaaaaatt ggaaaagggc 180 taaattatcc gtcggggtgt cctcctagct cggctcaacc taggcaaatg cgtttactgg 240 ggccatccaa gctcatcctt ccagagattc gcctttcaga ggcaaagaac tcgtctccgc 300 aggeetettg tteegggagg aggagaatte ttgegeggaa aetggttgat geetgggeta 360 tggtaattct gacacctttg ctatcctaac tggaaaaggt ccttagtaat aacaatatca 420 ggtactttaa cattgttaaa gatatactat tagtgaaaaa ccgatcttac gaaaagccaa 480 tagagcagaa agtggtaaag atgtctttag ttgtacaaga acaaggttcc ttccaacaca 540 ttttacggta tgttttacga gtacagtgca gatatgttga gagagtttca cattacatgg 600 gataatagct taatattgcg attgaaaata tgtgagtgca ctccgatgga ataataaacg 660 tcacgggtta ccaaatacga ggccgaatta actctacgct actataaaaa ttatcactag 720 attttaaagt aatacacagt ttattggaaa atacaatata tgactcaatg cgaatttaaa 780 aatttcagtc aatacagctt gacaaattca tatttatcgt ccctccatgt tcgtacgcca 840 ttccaaaatg atgcgccatg ttgaaaattt ctcgaagaac ctagcagtgg tgtgaaataa 900 attactaaca ataagatgta caattttttg tgcgcccatt acaatcagtt tgttgaacac 960 caatgtcgat ggtaacatta aaattgttta cgctttgacc actattaagg gtgttggtcg 1020 tegttactee aacttggtet gtaagaagge tgatgttgat ttgcacaaga gagetggtga 1080 attgacccaa gaagaattgg aaagaattgt ccaaatcatg caaaacccaa ctcattacaa 1140 gatcccagcc tggttcttaa accgtcaaaa cgacattact gatggtaagg actaccacac 1200 tttggctaac aacgtcgaat ccaagttgag agatgacttg gaaagattaa agaagatcag 1260 ageceacegt ggtattagae acttetgggg tttacgtgtt agaggteaae acaecaagae 1320 cactggtaga agaagagctt aa 1342

```
<210> 208
```

<211> 146

<212> PRT

<213> Saccharomyces cerevisiae

<400> 208

Met Ser Leu Val Val Gln Glu Gln Gly Ser Phe Gln His Ile Leu Arg
1 5 10 15

Leu Leu Asn Thr Asn Val Asp Gly Asn Ile Lys Ile Val Tyr Ala Leu 20 25 30

Thr Thr Ile Lys Gly Val Gly Arg Arg Tyr Ser Asn Leu Val Cys Lys
35 40 45

Lys Ala Asp Val Asp Leu His Lys Arg Ala Gly Glu Leu Thr Gln Glu 50 55 60

Glu Leu Glu Arg Ile Val Gln Ile Met Gln Asn Pro Thr His Tyr Lys
65 70 75 80

Ile Pro Ala Trp Phe Leu Asn Arg Gln Asn Asp Ile Thr Asp Gly Lys
85 90 95

Asp Tyr His Thr Leu Ala Asn Asn Val Glu Ser Lys Leu Arg Asp Asp 100 105 110

Leu Glu Arg Leu Lys Lys Ile Arg Ala His Arg Gly Ile Arg His Phe 115 120 125

Trp Gly Leu Arg Val Arg Gly Gln His Thr Lys Thr Thr Gly Arg Arg 130 135 140

Arg Ala

145

<210> 209

<211> 1268

<212> DNA

<213> Saccharomyces cerevisiae

<400> 209

tatattattt tttcccttct gggttctttt cttccttttc ttgtttacct tttttgcttt 60 ttcataaaat aatttctcta gatttgaaga cagcattttt gtacatccat acaccataca 120

```
ccatacacca tagcaccagt acactatatt tttatgaatt ttactaagaa ttattcctgc 180
aggageteca etgaaaaaa aagageagea tggatgteeg gtagaagtge taetgagtaa 240
atgggaagga cgcggtagat ccagtgtgga atcaaggtgg tgccggtgtg aagccgcctc 300
ggccggctgg actctccagg ccggagtgat gattgccacg ctgaacgtaa cacagtttca 360
caataccagt gtcctcatta gtgagttcca atgtatagtt agtagtggta ttttgatata 420
tgtgagtggt agcagatttg aacttagtta gttgtattcg cctttgagga aaccaagcaa 480
aaaaccgatc tagactaatc atggctgttg gtaagaataa gagactatcc agaggtaaga 540
agggtttgaa gaagaaggtc gttgacccat ttaccagaaa ggaatggttc gatattaaag 600
ccccatccac ttttgaaaac agaaatgttg gtaagacttt agttaacaag tccactggtt 660
tgaagaatgc ttccgatgct ttgaagggta gagttgtcga agtttgtttg gctgacttgc 720
aaggttetga agaccattet tteagaaagg teaagttgag agtegatgaa gtteaaggta 780
agaacttgtt gaccaacttc cacggtatgg acttcactac cgacaaattg agatcaatgg 840
tcagaaaatg gcaaactttg atcgaagcta atgttaccgt taagacttcc gatgattacg 900
ttttgagaat ctttgctatt gccttcacca gaaagcaagc taaccaagtt aagagacact 960
cttacgctca atcttcccac atcagagcta tcagaaaagt tatttctgaa atcttgacca 1020
gagaagttca aaactctact ttggctcaat tgacctccaa attgattcca gaagttatca 1080
acaaggaaat cgaaaatgct accaaggaca tcttcccact acaaaacatc cacgttagaa 1140
aggttaagtt attgaaacaa ccaaagttcg acgttggtgc tttgatggct ttgcatggtg 1200
aaggttccgg tgaagaaaag ggtaagaagg tttctggttt caaggatgaa gtcttggaaa 1260
                                                                  1268
ctgtgtaa
<210> 210
<211> 255
<212> PRT
```

<213> Saccharomyces cerevisiae

<400> 210

Met Ala Val Gly Lys Asn Lys Arg Leu Ser Arg Gly Lys Lys Gly Leu 10

Lys Lys Val Val Asp Pro Phe Thr Arg Lys Glu Trp Phe Asp Ile 20 25

Lys Ala Pro Ser Thr Phe Glu Asn Arg Asn Val Gly Lys Thr Leu Val 40

Asn Lys Ser Thr Gly Leu Lys Asn Ala Ser Asp Ala Leu Lys Gly Arg 50 55 60

Val Val Glu Val Cys Leu Ala Asp Leu Gln Gly Ser Glu Asp His Ser 65 70 75

Phe Arg Lys Val Lys Leu Arg Val Asp Glu Val Gln Gly Lys Asn Leu 85 90

Leu Thr Asn Phe His Gly Met Asp Phe Thr Thr Asp Lys Leu Arg Ser 100 105

SEQL

```
Met Val Arg Lys Trp Gln Thr Leu Ile Glu Ala Asn Val Thr Val Lys
        115
                            120
                                                 125
Thr Ser Asp Asp Tyr Val Leu Arg Ile Phe Ala Ile Ala Phe Thr Arg
                         135
                                             140
Lys Gln Ala Asn Gln Val Lys Arg His Ser Tyr Ala Gln Ser Ser His
145
                    150
                                        155
Ile Arg Ala Ile Arg Lys Val Ile Ser Glu Ile Leu Thr Arg Glu Val
                165
                                     170
Gln Asn Ser Thr Leu Ala Gln Leu Thr Ser Lys Leu Ile Pro Glu Val
            180
                                 185
Ile Asn Lys Glu Ile Glu Asn Ala Thr Lys Asp Ile Phe Pro Leu Gln
        195
                            200
                                                 205
Asn Ile His Val Arg Lys Val Lys Leu Leu Lys Gln Pro Lys Phe Asp
    210
                        215
                                            220
Val Gly Ala Leu Met Ala Leu His Gly Glu Gly Ser Gly Glu Glu Lys
225
                    230
                                        235
                                                             240
Gly Lys Lys Val Ser Gly Phe Lys Asp Glu Val Leu Glu Thr Val
                245
                                     250
<210> 211
<211> 2042
<212> DNA
<213> Saccharomyces cerevisiae
<400> 211
ttaatagggt tagccaccat tgtctcgaac atgttcacag gctcggagct tgtggcgggt 60
gcctcgctta ctggagtacc atttagcggt gtgtttgaag ctaacgagtt catcttcaat 120
ctttatttcc acttctttac aagctttctg cctttctcta tgacgtcttt cccaaaacac 180
tacctcttga cacgacttgt tgttttcgtt tttctctaag aatatcacta ttttcacttt 240
```

ccattgagga cttgaaaacg gaggcatcgc aggtatggga caaacacgcg cagcccaaac 720

tttcaccttt tccaccatta cgtcggtaag caacagcatt acgccggcct caccttagag 300 gggaagaata agggatcttt ggaactgaag gaaaataagg gagagggagg aaacaaggag 360 gcggtgaaag gggatcgaca gggtatttaa agccactcat ggagtcgtat tcctagcatg 420 attagattac cggagtttag aagaggatct gcaatagaag aggagacaac tacaaaagaa 480 ctgttaatta agagaagaag atgaagcaat tcaagttggt taatgcggtt tccgcatcat 540 ttgtgcttat tggcttagtg ttggccaatt cagattcagt gttcgacaag tggacgcagg 600 aagacctggc cgattatta cgtgataata agaagagctt ggaaaagtac gctacagact 660



```
catggtggca ggtgtggtct agtgacagca gtagcgtgag caacagtaac cccggatggt 780
ttggttatac tggttcttcg gatcacccag tttctgactg gctctttgac acctggtcca 840
cagacagtet acgtaactte ttgaagaaaa acggcgtgga tgttgacgac gctaaggett 900
ccaaggactc gctggtgaag actgctaagg agaacttcaa caagatttcc aaatctttga 960
agtcgtcggg atactatcct tctagctctt actttgatag ctggtcaacc aaagatttgc 1020
aaaactggtt gaatgacaat ggtatcgact acgacaaggc agttcaaagc aaggacgagc 1080
tagttcagaa agtcaaggaa aacatctacc gaacttcaga aaaggcagaa cagcagcgtc 1140
tgggtttgct agaaagcttg gatttggctc accaacaaat attagacaca tcgggacaaa 1200
taaaagacac tgtatttgac aagtggtcta gtgatcagtt aaccaattgg ttggagagcc 1260
acaaggtcaa tattgacaag aacatggcca agaaacacga ctatttggtt agaatggcca 1320
aagaaaattc tgccaatttg aaagatgata tctactggta cctggactat atgaaaagag 1380
agtettetee atttttgace aagaceeeag aataegttgg tteegtttgg gactetteta 1440
aaaatttcct cacaaatttg tactccaagt tcagaggtaa gactgacaat gtgatcaatg 1500
atactttttt ggttggccta gactcttggc caaaggacaa attgaaaatg tttttagatg 1560
ctcgtggtat taagtactca atgctgtcta ctgaacatca attgagagaa ttagtcaaaa 1620
aatccagaaa cgaaaaactc aagatcctgc caaaagacta ccaaaaatac tttgacaaca 1680
gtaactggtc tttggatgac ataaagggtt ggtttgctga caaaaaggac gacttccaag 1740
attctcagac ttactccaca attatgcagg attttgacaa ggtttccaaa aacacaaatg 1800
atgctaagga ccaaatcgct aagacctggt caaatacctt tcagagctgg tctcaagaag 1860
acctattgca gtacctaaaa tcattcggtg ttccggttaa acagacttct acgaaggacg 1920
acttaatcaa cttggccaag cagaatacgc aatggttgtt cggcactgtt aaggagcctg 1980
cttacaagag gtacctacat aacgttaaaa actggtcgaa aagcatatta gggttcaact 2040
aa
```

```
<210> 212
```

<211> 513

<212> PRT

<213> Saccharomyces cerevisiae

<400> 212

```
Met Lys Gln Phe Lys Leu Val Asn Ala Val Ser Ala Ser Phe Val Leu
1 5 10 15
```

Ile Gly Leu Val Leu Ala Asn Ser Asp Ser Val Phe Asp Lys Trp Thr 20 25 30

Gln Glu Asp Leu Ala Asp Tyr Leu Arg Asp Asn Lys Lys Ser Leu Glu
35 40 45

Lys Tyr Ala Thr Asp Ser Ile Glu Asp Leu Lys Thr Glu Ala Ser Gln 50 55 60

Val Trp Asp Lys His Ala Gln Pro Lys Pro Trp Trp Gln Val Trp Ser
65 70 75 80

Ser Asp Ser Ser Ser Val Ser Asn Ser Asn Pro Gly Trp Phe Gly Tyr 85 90 95

Thr	Gly	Ser	Ser 100	Asp	His	Pro	Val	Ser 105	Asp	Trp	Leu	Phe	Asp 110	Thr	Trp
Ser	Thr	Asp 115	Ser	Leu	Arg	Asn	Phe 120	Leu	Lys	Lys	Asn	Gly 125	Val	Asp	Val
Asp	Asp 130	Ala	Lys	Ala	Ser	Lys 135	Asp	Ser	Leu	Val	Lys 140	Thr	Ala	Lys	Glu
Asn 145	Phe	Asn	Lys	Ile	Ser 150	Lys	Ser	Leu	Lys	Ser 155	Ser	Gly	Туг	Туг	Pro
Ser	Ser	Ser	Tyr	Phe 165	Asp	Ser	Trp	Ser	Thr 170	Lys	Asp	Leu	Gln	Asn 175	Trp
Leu	Asn	Asp	Asn 180	Gly	Ile	Asp	Tyr	Asp 185	Lys	Ala	Val	Gln	Ser 190	Lys	Asp
Glu	Leu	Val 195	Gln	Lys	Val	Lys	Glu 200	Asn	Ile	Tyr	Arg	Thr 205	Ser	<b>Gl</b> u	Lys
Ala	Glu 210	Gln	Gln	Arg	Leu	Gly 215	Leu	Leu	Glu	Ser	Leu 220	Asp	Leu	Ala	His
Gln 225	Gln	Ile	Leu	Asp	Thr 230	Ser	Gly	Gln	Ile	Lys 235	Asp	Thr	Val	Phe	Asp 240
Lys	Trp	Ser	Ser	Asp 245	Gln	Leu	Thr	Asn	Trp 250	Leu	Glu	Ser	His	Lys 255	Val
Asn	Ile	Asp	Lys 260	Asn	Met	Ala	Lys	Lys 265	His	Asp	Tyr	Leu	Val 270	Arg	Met

Ala Lys Glu Asn Ser Ala Asn Leu Lys Asp Asp Ile Tyr Trp Tyr Leu 275 280 285

Asp Tyr Met Lys Arg Glu Ser Ser Pro Phe Leu Thr Lys Thr Pro Glu 290 295 300

Tyr Val Gly Ser Val Trp Asp Ser Ser Lys Asn Phe Leu Thr Asn Leu 305 310 315 320

Tyr Ser Lys Phe Arg Gly Lys Thr Asp Asn Val Ile Asn Asp Thr Phe 325 330 335

Leu Val Gly Leu Asp Ser Trp Pro Lys Asp Lys Leu Lys Met Phe Leu 340 345 350

```
Asp Ala Arg Gly Ile Lys Tyr Ser Met Leu Ser Thr Glu His Gln Leu 355 360 365
```

Arg Glu Leu Val Lys Lys Ser Arg Asn Glu Lys Leu Lys Ile Leu Pro 370 375 380

Lys Asp Tyr Gln Lys Tyr Phe Asp Asn Ser Asn Trp Ser Leu Asp Asp 385 390 395 400

Ile Lys Gly Trp Phe Ala Asp Lys Lys Asp Asp Phe Gln Asp Ser Gln 405 410 415

Thr Tyr Ser Thr Ile Met Gln Asp Phe Asp Lys Val Ser Lys Asn Thr 420 425 430

Asn Asp Ala Lys Asp Gln Ile Ala Lys Thr Trp Ser Asn Thr Phe Gln 435 440 445

Ser Trp Ser Gin Glu Asp Leu Cln Tyr Leu Lys Ser Phe Gly Val 450 455 460

Pro Val Lys Gln Thr Ser Thr Lys Asp Asp Leu Ile Asn Leu Ala Lys 465 470 475 480

Gln Asn Thr Gln Trp Leu Phe Gly Thr Val Lys Glu Pro Ala Tyr Lys 485 490 495

Arg Tyr Leu His Asn Val Lys Asn Trp Ser Lys Ser Ile Leu Gly Phe 500 505 510

Asn

<210> 213

<211> 2192

<212> DNA

<213> Saccharomyces cerevisiae

<400> 213

acgagatcat tttcttatct atctattgag taatgcttac ttttcatatt ttcaatgaac 60 aataggatat gtaggagaat tgatatattc actgcgtatc agagaaaagg tctactgaca 120 ttttatggca aatgtattct acacaaatcg agaataccac agacaatggt acaagacata 180 cacaaagaga agactgttct aattaaacaa ataatattga gctacctgct aagtatgtcc 240 ttttcccttt gtcctttggt ttctcttata gaagaccctg gaaatttttc gcatttttcc 300 ggctttgggc gttagtaaga acaaaaagaa aagaagagaa caaaaaagaa acgatacgga 360



```
gtacgtgtca taaaaacttg ttcaatcatc cttgaagcta agtataaaga gcttgaaaag 420
gtttaccact taaactggtt atactatttc aagagtgtaa acattttatt gcatatacca 480
cagtaacgtg caggtaaaac atgagattaa gaaccgccat tgccacactg tgcctcacgg 540
cttttacatc tgcaacttca aacaatagct acatcgccac cgaccaaaca caaaatgcct 600
ttaatgacac tcacttttgt aaggtcgaca ggaatgatca cgttagtccc agttgtaacg 660
taacattcaa tgaattaaat gccataaatg aaaacattag agatgatctt tcggcgttat 720
taaaatctga tttcttcaaa tactttcggc tggatttata caagcaatgt tcattttggg 780
acgccaacga tggtctgtgc ttaaaccgcg cttgctctgt tgatgtcgta gaggactggg 840
atacactgcc tgagtactgg cagcctgaga tcttgggtag tttcaataat gatacaatga 900
aggaagcgga tgatagcgat gacgaatgta agttcttaga tcaactatgt caaaccagta 960
aaaaacctgt agatatcgaa gacaccatca actactgtga tgtaaatgac tttaacggta 1020
aaaacgccgt tctgattgat ttaacagcaa atccggaacg atttacaggt tatggtggta 1080
agcaagctgg tcaaatttgg tctactatct accaagacaa ctgttttaca attggcgaaa 1140
ctggtgaatc attggccaaa gatgcatttt atagacttgt atccggtttc catgcctcta 1200
tcggtactca cttatcaaag gaatatttga acacgaaaac tggtaaatgg gagcccaatc 1260
tggatttgtt tatggcaaga atcgggaact ttcctgatag agtgacaaac atgtatttca 1320
attatgctgt tgtagctaag gctctctgga aaattcaacc atatttacca gaattttcat 1380
tctgtgatct agtcaataaa gaaatcaaaa acaaaatgga taacgttatt tcccagctgg 1440
acacaaaat ttttaacgaa gacttagttt ttgccaacga cctaagtttg actttgaagg 1500
acgaattcag atctcgcttc aagaatgtca cgaagattat ggattgtgtg caatgtgata 1560
gatgtagatt gtggggcaaa attcaaacta ccggttacgc aactgccttg aaaattttgt 1620
ttgaaatcaa cgacgctgat gaattcacca aacaacatat tgttggtaag ttaaccaaat 1680
atgagttgat tgcactatta cagactttcg gtagattatc tgaatctatt gaatctgtta 1740
acatgttcga aaaaatgtac gggaaaaggt taaacggttc tgaaaacagg ttaagctcat 1800
tcttccaaaa taacttcttc aacattttga aggaggcagg caaatcgatt cgttacacca 1860
tagagaacat caattccact aaagaaggaa agaaaaagac taacaattct caatcacatg 1920
tatttgatga tttaaaaatg cccaaagcag aaatagttcc aaggccctct aacggtacag 1980
taaataaatg gaagaaagct tggaatactg aagttaacaa cgttttagaa gcattcagat 2040
ttatttatag aagctatttg gatttaccca ggaacatctg ggaattatct ttgatgaagg 2100
tatacaaatt ttggaataaa ttcatcggtg ttgctgatta cgttagtgag gagacacgag 2160
                                                                  2192
agcctatttc ctataagcta gatatacaat aa
```

```
<210> 214
```

<211> 563

<212> PRT

<213> Saccharomyces cerevisiae

## <400> 214

Met Arg Leu Arg Thr Ala Ile Ala Thr Leu Cys Leu Thr Ala Phe Thr 1 5 10 15

Ser Ala Thr Ser Asn Asn Ser Tyr Ile Ala Thr Asp Gln Thr Gln Asn 20 25 30

Ala Phe Asn Asp Thr His Phe Cys Lys Val Asp Arg Asn Asp His Val
35 40 45



- Ser Pro Ser Cys Asn Val Thr Phe Asn Glu Leu Asn Ala Ile Asn Glu
  50 55 60
- Asn Ile Arg Asp Asp Leu Ser Ala Leu Leu Lys Ser Asp Phe Phe Lys 65 70 75 80
- Tyr Phe Arg Leu Asp Leu Tyr Lys Gln Cys Ser Phe Trp Asp Ala Asn 85 90 95
- Asp Gly Leu Cys Leu Asn Arg Ala Cys Ser Val Asp Val Val Glu Asp 100 105 110
- Trp Asp Thr Leu Pro Glu Tyr Trp Gln Pro Glu Ile Leu Gly Ser Phe
  115 120 125
- Asn Asn Asp Thr Met Lys Glu Ala Asp Asp Ser Asp Asp Glu Cys Lys
  130 135 140
- Phe Leu Asp Gln Leu Cys Gln Thr Ser Lys Lys Pro Val Asp Ile Glu 145 150 155 160
- Asp Thr Ile Asn Tyr Cys Asp Val Asn Asp Phe Asn Gly Lys Asn Ala 165 170 175
- Val Leu Ile Asp Leu Thr Ala Asn Pro Glu Arg Phe Thr Gly Tyr Gly
  180 185 190
- Gly Lys Gln Ala Gly Gln Ile Trp Ser Thr Ile Tyr Gln Asp Asn Cys 195 200 205
- Phe Thr Ile Gly Glu Thr Gly Glu Ser Leu Ala Lys Asp Ala Phe Tyr 210 215 220
- Arg Leu Val Ser Gly Phe His Ala Ser Ile Gly Thr His Leu Ser Lys 225 230 235 240
- Glu Tyr Leu Asn Thr Lys Thr Gly Lys Trp Glu Pro Asn Leu Asp Leu 245 250 255
- Phe Met Ala Arg Ile Gly Asn Phe Pro Asp Arg Val Thr Asn Met Tyr 260 265 270
- Phe Asn Tyr Ala Val Val Ala Lys Ala Leu Trp Lys Ile Gln Pro Tyr 275 280 285
- Leu Pro Glu Phe Ser Phe Cys Asp Leu Val Asn Lys Glu Ile Lys Asn 290 295 300

Lys	Met	Asp	Asn	Val	Ile	Ser	Gln	Leu	Asp	Thr	Lys	Ile	Phe	Asn	Glu
305					310					315					320

Asp Leu Val Phe Ala Asn Asp Leu Ser Leu Thr Leu Lys Asp Glu Phe 325 330 335

Arg Ser Arg Phe Lys Asn Val Thr Lys Ile Met Asp Cys Val Gln Cys 340 345 350

Asp Arg Cys Arg Leu Trp Gly Lys Ile Gln Thr Thr Gly Tyr Ala Thr
355 360 365

Ala Leu Lys Ile Leu Phe Glu Ile Asn Asp Ala Asp Glu Phe Thr Lys 370 375 380

Gln His Ile Val Gly Lys Leu Thr Lys Tyr Glu Leu Ile Ala Leu Leu 385 390 395 400

Gln Thr Phe Gly Arg Leu Ser Glu Ser Ile Glu Ser Val Asn Met Phe 405 410 415

Glu Lys Met Tyr Gly Lys Arg Leu Asn Gly Ser Glu Asn Arg Leu Ser 420 425 430

Ser Phe Phe Gln Asn Asn Phe Phe Asn Ile Leu Lys Glu Ala Gly Lys
435
440
445

Ser Ile Arg Tyr Thr Ile Glu Asn Ile Asn Ser Thr Lys Glu Gly Lys 450 455 460

Lys Lys Thr Asn Asn Ser Gln Ser His Val Phe Asp Asp Leu Lys Met 465 470 475 480

Pro Lys Ala Glu Ile Val Pro Arg Pro Ser Asn Gly Thr Val Asn Lys
485 490 495

Trp Lys Lys Ala Trp Asn Thr Glu Val Asn Asn Val Leu Glu Ala Phe
500 505 510

Arg Phe Ile Tyr Arg Ser Tyr Leu Asp Leu Pro Arg Asn Ile Trp Glu 515 520 525

Leu Ser Leu Met Lys Val Tyr Lys Phe Trp Asn Lys Phe Ile Gly Val 530 535 540

Ala Asp Tyr Val Ser Glu Glu Thr Arg Glu Pro Ile Ser Tyr Lys Leu 545 550 555 560

Asp Ile Gln

<210> 215 <211> 998 <212> DNA <213> Saccharomyces cerevisiae

<400> 215

aacagtattg gctgttgatt cattgcgctg cagcagtcac cttcaatttg tgcaccattt 60 tcgtattctg tactttgcga tgtagagtct actaaaatag cgtctctgat agccatgggt 120 gaaggtette etetagttet cacettaatt agcatteggt gagaatgeet geatgttgaa 180 gagcgatgcc ctctgatgca cgatgcacac gcatatttgt tcccattaaa tattatcatc 240 tctgatagag cttgaagaac ttaccagact gtttcaagtt taaaacaagg cgcctcatcg 300 catctacctc acgaagatgc agcagcttat tgtctcgaga tccttcttat aactcctttt 360 cgccattacc cgaaaacgag attacagcct ctaaaccaag agctcgaaaa gcgccaaagt 420 aaactctggc gtttagcgta cgaaggagat tatcctaaaa ggaacttccc tagtaatagt 480 gtaatttgga agggcatage atgtegaaaa eegeteagaa aegteteete aaggagette 540 aacagttaat taaagattct ccacctggta tagtggctgg tcccaaatcg gagaataaca 600 tattcatttg ggactgccta attcaagggc ctccagatac gccatacgct gatggtgttt 660 ttaatgctaa gctagagttt cctaaagact atccgttatc tccacctaaa cttactttca 720 cacccagcat actacatcca aatatttatc caaatgggga agtgtgcata tccattctac 780 actcccctgg tgatgatcct aacatgtacg aattagcgga agaaagatgg tcgccagtgc 840 aaagtgtaga aaaaatteta ttaagtgtta tgagcatgtt gagtgageee aatategaaa 900 gtggtgccaa cattgatgct tgcatcttgt ggagagataa tagacctgaa tttgagagac 960 998 aggtaaagtt atccattttg aaatcattag gattctga

<210> 216 <211> 165 <212> PRT <213> Saccharomyces cerevisiae

<400> 216

Met Ser Lys Thr Ala Gln Lys Arg Leu Leu Lys Glu Leu Gln Gln Leu
1 5 10 15

Ile Lys Asp Ser Pro Pro Gly Ile Val Ala Gly Pro Lys Ser Glu Asn 20 25 30

Asn Ile Phe Ile Trp Asp Cys Leu Ile Gln Gly Pro Pro Asp Thr Pro
35 40 45

Tyr Ala Asp Gly Val Phe Asn Ala Lys Leu Glu Phe Pro Lys Asp Tyr 50 55 60

Pro Leu Ser Pro Pro Lys Leu Thr Phe Thr Pro Ser Ile Leu His Pro
65 70 75 80

Asn Ile Tyr Pro Asn Gly Glu Val Cys Ile Ser Ile Leu His Ser Pro

Asn Ile Tyr Pro Asn Gly Glu Val Cys Ile Ser Ile Leu His Ser Pro 85 90 95

Gly Asp Asp Pro Asn Met Tyr Glu Leu Ala Glu Glu Arg Trp Ser Pro 100 105 110

Val Gln Ser Val Glu Lys Ile Leu Leu Ser Val Met Ser Met Leu Ser 115 120 125

Glu Pro Asn Ile Glu Ser Gly Ala Asn Ile Asp Ala Cys Ile Leu Trp 130 135 140

Arg Asp Asn Arg Pro Glu Phe Glu Arg Gln Val Lys Leu Ser Ile Leu 145 150 155 160

Lys Ser Leu Gly Phe 165

<210> 217

<211> 1091

<212> DNA

<213> Saccharomyces cerevisiae

## <400> 217

gtccttccgc tttatatgtt tcgttatcct atcacattat caaatgactt tgtgccactg 60 tgttttgaca aaattactgt aatataatca atagtattta catgtttgta ccagaagtac 120 ttgcatcgca tccagttctc atgcaatata gttgtatacc atgtcgttga aaggaaccag 180 agtaaacact tctaccagta tttctttacg gttcggatca aaaccatcac tcattcggtc 240 attettaceg tacgataaac caaattacaa ggaaaagtat ceetaattag agateaetge 300 atccgtaata gtttttttc tattttggac ttttgtaaaa aagggattag ggatacgttg 360 ctcataaaaa aattgacgaa gattttagat aatggcaaat aaaaatgaaa tagtatcaat 420 ataccgaaaa attaatcaca ctcaatgcga ctgtgatagc tgataagtgg agctcagaaa 480 tattcagaag cgtaagaata atgaaagcaa ccattcaaag agtaacatct gtatttggag 540 ttccccgagc atctgtattc gtgccaagaa tcagcacacc atttattttg cataattata 600 tetetaatgg cagaatggae ettttteea aagaatteea caatggeege gtateeaaat 660 ctgatctttg gtcaagcaac aaggaagagg agctcttagt atctcaaagg aaaaaaagac 720 egatetetee teatttaact gtttacgaac etgaaatgag etggtatett teetetttac 780 atcgtatatc tggcgtttta ctggctcttg gattctatgc tttcacaatt actttgggtg 840 tgacgacaat aatgggaatg gatacgactt ttcaggatct aaacaagtgg tatcacgaaa 900 agatgcctaa gtggtcccaa tgggtggcca aaggctctgc agcatatctg tttgcattcc 960 attttggtaa cggtataagg catctcatct gggatatggg ctacgaattg accaaccgtg 1020 gggttataaa aaccggatca atcgttttag ccggcacact cgtcttagga acgtatttac 1080 tggctcagta a

<210> 218

<211> 196

<212> PRT

<213> Saccharomyces cerevisiae

<400> 218

Met Lys Ala Thr Ile Gln Arg Val Thr Ser Val Phe Gly Val Pro Arg

1 5 10 15

Ala Ser Val Phe Val Pro Arg Ile Ser Thr Pro Phe Ile Leu His Asn 20 25 30

Tyr Ile Ser Asn Gly Arg Met Asp Leu Phe Ser Lys Glu Phe His Asn 35 40 45

Gly Arg Val Ser Lys Ser Asp Leu Trp Ser Ser Asn Lys Glu Glu Glu 50 55 60

Leu Leu Val Ser Gln Arg Lys Lys Arg Pro Ile Ser Pro His Leu Thr
65 70 75 80

Val Tyr Glu Pro Glu Met Ser Trp Tyr Leu Ser Ser Leu His Arg Ile 85 90 95

Ser Gly Val Leu Leu Ala Leu Gly Phe Tyr Ala Phe Thr Ile Thr Leu 100 105 110

Gly Val Thr Thr Ile Met Gly Met Asp Thr Thr Phe Gln Asp Leu Asn 115 120 125

Lys Trp Tyr His Glu Lys Met Pro Lys Trp Ser Gln Trp Val Ala Lys 130 135 140

Gly Ser Ala Ala Tyr Leu Phe Ala Phe His Phe Gly Asn Gly Ile Arg 145 150 155 160

His Leu Ile Trp Asp Met Gly Tyr Glu Leu Thr Asn Arg Gly Val Ile 165 170 175

Lys Thr Gly Ser Ile Val Leu Ala Gly Thr Leu Val Leu Gly Thr Tyr 180 185 190

Leu Leu Ala Gln 195 <210> 219 <211> 1121

```
<212> DNA
<213> Saccharomyces cerevisiae
<400> 219
ccgcttagcg caaactatcg tgaactcgct gcaacaactg agagggcaag gatatacata 60
aaaatagcct acaaattctg aactctgtaa aggaagcctc ataaataaag gtagatagta 120
aagtatacaa gagaagaatc ccaagatgtc agctgtccca agtgttcaag tatgttttca 180
gttctgcaga atgatgtttg atagtatcga taatggagtg agatcaagag aaaaaaaatg 240
aatatgtcag ccaaccaagt tctgagtagg cagtaaatga gtacgcatag tgtatttatc 300
caaaggaaag aattgttatt tttacaagcc gaattgagat ccaattaggc aatgttttgg 360
ggagagtatt ttgacaagat tggttaaact actacggtca gttccgtaac cagtacgatt 420
gtacacataa ggaaacaact gtaaagataa acaataaggg cttccaatgc cattgtaaga 480
tatcatattc ctaaacaaaa atgtacagcg aatataaagc cagcgtcagt gtcttcctgg 540
aaggttgcgc aactaaagaa gttgaaatta aaatggcaag cgctacattt tcatccattt 600
tcactcatca gatgtccaag attttcgaaa aataaatatt ctcatttttc tctcaatgaa 660
ataattgtta ctaacattga atttcctcgt aactaattgc attacttctt tagacttttg 720
gtaagaagaa atcagctact gctgttgccc atgtcaaggc cggtaagggt ttgatcaagg 780
ttaatggttc tccaatcact ttggttgaac cagaaatctt aagattcaag gtttacgaac 840
cattattgtt agttggtttg gacaaattct ccaacatcga tattagagtt agagttactg 900
gtggtggtca tgtttcccaa gtttacgcca tcagacaagc tattgctaaa ggtttagttg 960
cttaccatca aaaatatgtc gatgaacaat ccaagaacga attgaagaag gctttcactt 1020
cttacgacag aaccttgttg attgctgatt ctagaagacc agaaccaaag aaattcggtg 1080
gtaagggtgc tcgttccaga ttccaaaaat cttaccgtta a
                                                                  1121
<210> 220
<211> 143
<212> PRT
<213> Saccharomyces cerevisiae
Met Tyr Ser Glu Tyr Lys Ala Ser Thr Phe Gly Lys Lys Lys Ser Ala
 1
                  5
                                     10
                                                          15
Thr Ala Val Ala His Val Lys Ala Gly Lys Gly Leu Ile Lys Val Asn
             20
                                 25
Gly Ser Pro Ile Thr Leu Val Glu Pro Glu Ile Leu Arg Phe Lys Val
         35
                             40
                                                 45
Tyr Glu Pro Leu Leu Val Gly Leu Asp Lys Phe Ser Asn Ile Asp
                         55
Ile Arg Val Arg Val Thr Gly Gly Gly His Val Ser Gln Val Tyr Ala
```

```
Ile Arg Gln Ala Ile Ala Lys Gly Leu Val Ala Tyr His Gln Lys Tyr
Val Asp Glu Gln Ser Lys Asn Glu Leu Lys Lys Ala Phe Thr Ser Tyr
                                105
            100
Asp Arg Thr Leu Leu Ile Ala Asp Ser Arg Arg Pro Glu Pro Lys Lys
                            120
Phe Gly Gly Lys Gly Ala Arg Ser Arg Phe Gln Lys Ser Tyr Arg
    130
                        135
<210> 221
<211> 707
<212> DNA
<213> Saccharomyces cerevisiae
<400> 221
qtcccatcat tctttacacc tcgtactgta ttcatgatca tctatccatt ttacataccg 60
ctccctttaa gatacgttta ttcgtaactc ccatttacca atcactatga gccgtccacg 120
tttccaaaaa aacctaaaca tatgatgcaa acctccaatg agactcaacg taacatgcaa 180
gtaaatacag aaggttaaga gatagttgtc ttaaaggggt accgaaagca tttaggggag 240
gcttaaggga gggtgccgat cctatgaagt attaatacgt aatgccaaaa ggaattgttg 300
aacatctgaa gtggatagat taatcgtaca gtaatcgtac agtactatgt cttactgatg 360
tegggaatet eaggggegae egeeceegge tagaattate tatataaegg taaaaagaat 420
aaaactctat tctagttctc gccatttacc ttgactttaa tgaaccaata aaagaaattt 480
ctacaaccaa gacatccaga atgaatacag accaacaaaa agtgagcgaa atatttcaga 540
gctcaaagga aaaattgcag ggcgatgcaa aggtagtgag tgacgctttt aagaaaatgg 600
ctagtcaaga caaggacggc aagactaccg atgctgatga aagtgaaaaa cacaactatc 660
                                                                   707
aagagcaata caacaagctc aaaggggcgg ggcataagaa ggagtag
<210> 222
<211> 68
<212> PRT
<213> Saccharomyces cerevisiae
<400> 222
Met Asn Thr Asp Gln Gln Lys Val Ser Glu Ile Phe Gln Ser Ser Lys
                                     10
```

Glu Lys Leu Gln Gly Asp Ala Lys Val Val Ser Asp Ala Phe Lys Lys
20 25 30

Met Ala Ser Gln Asp Lys Asp Gly Lys Thr Thr Asp Ala Asp Glu Ser

35

40

45

Glu Lys His Asn Tyr Gln Glu Gln Tyr Asn Lys Leu Lys Gly Ala Gly
50 55 60

His Lys Lys Glu 65

<210> 223

<211> 1877

<212> DNA

<213> Saccharomyces cerevisiae

<400> 223

ggttcgatac aacacttgtg ctggctggta tatcaaccat gggtacgtca gcatattgat 60 tcgatgtggc agtgtacgag agccattctg tatcgttcga caaagtttgc atgacttcat 120 gttgtttcgt agcagtattc gttggtccag atgcaggaat gctggttata aagtttgttg 180 ttcctggtcc acattcatct ttggatggcg ttgttggtgg cgaatttgtc ttaggccttt 240 tgttcgcgat gtgcttttgg atagcctgga atacgaaatc tttgtcttcc tgtaaagtcg 300 ccgtttcctt taaccatttg tgctcccttc ttaagttatg tgacggcttc gtcaccatta 360 ccgccttgtg tgtacgtgta tgatttttta aatatataca acaataatct gtattttttc 420 ctttcctcta gccaatgact ccaagctggc tgataaaaac aaactaaacg gtaaagccac 480 aaatccgaaa tgtatcacca atgatcaccc agcctgctaa gtgccctcta ttgatccgta 540 tatcagcttt tagatcaggc tcgagtttct tgttatatgt gcattgcaaa agcataaaca 600 aatcctggca gccgaagccg ggcaatccac ttcgaaacgc acggctgaac tatataaata 660 taaaggacat gtggagagaa gcttctcttc cttcacattt cgcatttcat gatctaaagt 720 ggttctttca caatagaaga gcaccaacac gaaatatggc tgtcggtggt aataactgga 780 gcatgtggct gcgaatgtca cgggtccacc tcaggcagat cacgaagtct ctggaccgaa 840 cgctgatcag tttgagccat gggaattttt cccaccaata taatcgcaat atttttgtca 900 cttggtggaa gagcctattt gaggcttcta cagcttttag gagggcaagt ggtttaacgg 960 tateteetet eaegaggaga ggaategeaa ggtttgacea ttteagacet gtteecaatg 1020 tcagtaaatt tgcttctttt cccagagtgc ccaaaggagc cccaaggggc ctttttacca 1080 actggaatat gactacatca aagagattat tagggcagag agcttattcg acttccagta 1140 tcaaatttac ccaagaagcc gtgaataaca tgactatatc tttgaggtgt tttttcaact 1200 cactaggcgg attaaatcag tgctcccact ctaattcctg caaagcttat cagaatgcgt 1260 ctaatgttac ctctaaacag gatcacgtcc aaccagttgc ccttaagaag ttatctcaaa 1320 aggatatcaa tttcattcgt aatttagaac tatttaagat aatgaagacc cagaatgaag 1380 tcgttgatga aacaagcgca tattacatgg aaaaaccagg ttcctatatt gaatttacca 1440 tttcagaatt taacgttaat gggacattct ccgcaccttt gtcattttta gatccttctt 1500 tgttagcgga tttggatgaa atgattagaa attacaaata cgaattaaaa tcaatataca 1560 gtagtgttga catgatcttg cagaattatg gatcattgcc gataactttt catcgaaata 1620 agattcgaat acattttcca aactcaactg tggtggaaac agaaaaacta attgcaggtc 1680 tgaatattgc tacgggtgtt atttacgcag atacgtctcc cgatatcagt ctagaaggta 1740 caaatttgaa tgctctggtt aatgttgaca attcaggaag cgtatggtct tttgttaagg 1800 agecetegtt tecetetagg agegettttt cacetatttt ateagatgea tectatgata 1860 cttatgaatt ggtctag 1877

<210> 224

<211> 458

<212> PRT

<213> Saccharomyces cerevisiae

<400> 224

Met Ile Thr Gln Pro Ala Lys Cys Pro Leu Leu Ile Arg Ile Ser Ala 1 5 10 15

Phe Arg Ser Gly Ser Ser Phe Leu Leu Tyr Val His Cys Lys Ser Ile 20 25 30

Asn Lys Ser Trp Gln Pro Lys Pro Gly Asn Pro Leu Arg Asn Ala Arg 35 40 45

Leu Asn Tyr Ile Asn Ile Lys Asp Met Trp Arg Glu Ala Ser Leu Pro 50 55 60

Ser His Phe Ala Phe His Asp Leu Lys Trp Phe Phe His Asn Arg Arg 65 70 75 80

Ala Pro Thr Arg Asn Met Ala Val Gly Gly Asn Asn Trp Ser Met Trp 85 90 95

Leu Arg Met Ser Arg Val His Leu Arg Gln Ile Thr Lys Ser Leu Asp 100 105 110

Arg Thr Leu Ile Ser Leu Ser His Gly Asn Phe Ser His Gln Tyr Asn 115 120 125

Arg Asn Ile Phe Val Thr Trp Trp Lys Ser Leu Phe Glu Ala Ser Thr 130 135 140

Ala Phe Arg Arg Ala Ser Gly Leu Thr Val Ser Pro Leu Thr Arg Arg 145 150 155 160

Gly Ile Ala Arg Phe Asp His Phe Arg Pro Val Pro Asn Val Ser Lys 165 170 175

Phe Ala Ser Phe Pro Arg Val Pro Lys Gly Ala Pro Arg Gly Leu Phe 180 185 190

Thr Asn Trp Asn Met Thr Thr Ser Lys Arg Leu Leu Gly Gln Arg Ala 195 200 205

Tyr Ser Thr Ser Ser Ile Lys Phe Thr Gln Glu Ala Val Asn Asn Met

SEQL

210 215 220

Thr Ile Ser Leu Arg Cys Phe Phe Asn Ser Leu Gly Gly Leu Asn Gln 225 230 235 240

Cys Ser His Ser Asn Ser Cys Lys Ala Tyr Gln Asn Ala Ser Asn Val 245 250 255

Thr Ser Lys Gln Asp His Val Gln Pro Val Ala Leu Lys Lys Leu Ser 260 265 270

Gln Lys Asp Ile Asn Phe Ile Arg Asn Leu Glu Leu Phe Lys Ile Met 275 280 285

Lys Thr Gln Asn Glu Val Val Asp Glu Thr Ser Ala Tyr Tyr Met Glu 290 295 300

Lys Pro Gly Ser Tyr Ile Glu Phe Thr Ile Ser Glu Phe Asn Val Asn 305 310 315 320

Gly Thr Phe Ser Ala Pro Leu Ser Phe Leu Asp Pro Ser Leu Leu Ala 325 330 335

Asp Leu Asp Glu Met Ile Arg Asn Tyr Lys Tyr Glu Leu Lys Ser Ile 340 345 350

Tyr Ser Ser Val Asp Met Ile Leu Gln Asn Tyr Gly Ser Leu Pro Ile 355 360 365

Thr Phe His Arg Asn Lys Ile Arg Ile His Phe Pro Asn Ser Thr Val 370 375 380

Val Glu Thr Glu Lys Leu Ile Ala Gly Leu Asn Ile Ala Thr Gly Val 385 390 395 400

Ile Tyr Ala Asp Thr Ser Pro Asp Ile Ser Leu Glu Gly Thr Asn Leu
405 410 415

Asn Ala Leu Val Asn Val Asp Asn Ser Gly Ser Val Trp Ser Phe Val
420 425 430

Lys Glu Pro Ser Phe Pro Ser Arg Ser Ala Phe Ser Pro Ile Leu Ser 435 440 445

Asp Ala Ser Tyr Asp Thr Tyr Glu Leu Val 450 455

```
<210> 225
<211> 1228
<212> DNA
<213> Saccharomyces cerevisiae
<400> 225
atcaaatatt gatcgagttc atataaagct aaagagtagg aaatatacca tcaatggtga 60
gcacttattt ttttttggat ttagatgtga gacatactaa aaaaaagttg ctatcaagcc 120
tataatttgg ctacgttgtc ttccggtgtt tttcaattga tttaagttac aacactcaaa 180
tctgggtaat ttgatctttt ttaataatta ttttagtgac atatagttct tagagttcgc 240
agatttattt tgtcattttt tggatcggcg tcttataaaa tcaagacata atacatccgc 300
acategegea tgtgtgggtg tatatgeeet ttteaegatt tttaagttge gteteaaaat 360
agtttccgag ttggaagcct gagtttttca aaacaatata ggaattaaag gtatacgtct 420
ttggatacat gttatttgaa atggggtaga actagcacaa ctgaaaccaa gaaaacacag 480
atcataacta accgttcaag atgttgatgc caaagcaaga aagaaacaaa attcaccaat 540
acttgtttca aggtatgttt gcatttttta ggtgaaatat gcagtgatat gctccgaaat 600
ggatagcaaa gatgataaat gaaaatacaa ttaaattgaa cttgaatatc attaaaagtg 660
gtagaaaacc gattgtttca aaaagatctt tgtggacaag tacggcagtg catgaatccg 720
aaagaaaagt gcaacctaag cagaattcca ttattcacg cgtccatttt tataatgttt 780
gaacttttga gtccttatag aaatgctaat attatgatca aagcgattat atcatttact 840
ttacagtttc cggagtttcc gtattattgc aaaaggacca caaacaattg aaatcatgat 900
atcqtaaqaa atattattac taacaaggaa gatcattgaa ttacqatcqc atatcqaaat 960
agaaggtgtt gtcgtcgcta agaaggattt caaccaagct aagcacgaag aaattgatac 1020
caagaacttg tatgttatca aggctttgca atccttgact tctaagggtt acgttaagac 1080
tcaattctca tggcaatact actattacac tttgactgaa gaaggtgttg aatacttgag 1140
agaatacttg aacttgcctg aacacattgt cccaggtacc tacattcaag aaagaaaccc 1200
                                                                  1228
atcccaaaga cctcaaagaa gatactaa
<210> 226
<211> 105
<212> PRT
<213> Saccharomyces cerevisiae
<400> 226
Met Leu Met Pro Lys Gln Glu Arg Asn Lys Ile His Gln Tyr Leu Phe
  1
                                     10
Gln Glu Gly Val Val Val Ala Lys Lys Asp Phe Asn Gln Ala Lys His
Glu Glu Ile Asp Thr Lys Asn Leu Tyr Val Ile Lys Ala Leu Gln Ser
                             40
```

Leu Thr Ser Lys Gly Tyr Val Lys Thr Gln Phe Ser Trp Gln Tyr Tyr

60

```
Tyr Tyr Thr Leu Thr Glu Glu Gly Val Glu Tyr Leu Arg Glu Tyr Leu 65 70 75 80
```

Asn Leu Pro Glu His Ile Val Pro Gly Thr Tyr Ile Gln Glu Arg Asn 85 90 95

Pro Ser Gln Arg Pro Gln Arg Arg Tyr
100 105

<210> 227 <211> 3998 <212> DNA <213> Saccharomyces cerevisiae

#### <400> 227

cccttcgcta aatcattaag aggtcattca tgaagttaat tcagcaccaa agttggactg 60 ttagacacga gatacacatt gcagtccaaa tgtgcagatt tttcagataa ttgtcgtcgc 120 acaagactgc ccctcccgtc ctcagacaga cacaaaatta gattttacgt ttacataaag 180 atgtacataa cttgaaaaaa cgttaatccc tcttaaattt agaccttgta agtcttcctg 240 gccactcttt gtgatatatt gtagtgcaat atattgattc gttcttctta tttctttgtt 300 cttaggcatt tccgctttag aaaattcgtt gggtggtttc tgcgacgggt atccccttcg 360 attttgcata atgatettea attetaeaae taaaateaag tagataeagg aaaatattee 420 ataaattata gtgtaaatcg ccctgtatac accttatcgt ttcatctcag gcaagttaaa 480 gcatttggga aacgtgctag atgacagaag aagatagaaa gctcactgta gagacagaaa 540 cagttgaggc accegtggca aataatettt tattgtcgaa taacagtaat gtagtagcac 600 ctaatcette tatteeetet geeteeacat etacetetee getacacagg gaaatagttg 660 atgattetgt egetaetget aacaceaeca geaaegttgt acageataat ttgeecaeca 720 tagataacaa tttaatggat teegatgeea egteacataa teaagateat tggeatteag 780 acataaacag ggcaggaaca tcaatgtcaa cgagtgatat cccaacagat ttacatttag 840 aacatatogg ototgtttoa toaactaata ataatagtaa caatgoocta atcaaccaca 900 accetetgte ateteatett tecaateegt catetteatt aegaaacaag aagagetett 960 tgttggtagc ttctaaccct gcgtttgctt cagatgttga actctcgaag aagaaacctg 1020 ccgtcatctc caataatatg cctacaagta acattgccct ttatcaaaca gcgagatcgg 1080 cgaatattca tggtccatca tcaacttccg catctaaagc gttcagaaag gcttcggcct 1140 tctccaataa cacggcaccc agcactagta ataacatcgg ctcgaataca cctccagctc 1200 ctcttttacc tctaccttca ctatcacaac aaaataagcc aaaaataata gagaggccca 1260 caatgcacgt cactaattca agagaaatac ttttaggtga aaacctgtta gatgatacaa 1320 aggogaagaa tgotocogog aattoaacoa cacaogataa tggtocagta gcaaatgatg 1380 ggctgcgtat accgaatcac tcgaacgcag atgataatga aaataacaac aaaatgaaga 1440 agaataaaaa cattaatagt gggaaaaatg aacgtaatga tgacaccagc aaaatatgca 1500 ctacatctac taaaacagcg ccttcaaccg cacctttggg cagtacagac aatactcagg 1560 ctctcactgc tagtgtctcc agcagtaatg ctgacaatca caataataac aagaagaaaa 1620 ccagcagcaa caacaacggc aataacagta atagtgcatc caataaaacc aatgccgata 1680 tcaagaattc taacgccgac ttgagcgctt ctacctctaa caataatgca ataaatgacg 1740 actcacatga gagtaattca gaaaaaccaa caaaggcgga ttttttcgct gcaaggctgg 1800 ctacagctgt aggtgaaaat gaaattagtg attctgagga aacatttgtt tatgaatcgg 1860

```
cagctaattc gactaaaaac ctaatatttc ctgactcctc cagccagcag cagcagcagc 1920
aacagcaacc tccaaaacaa cagcaacagc aacaaaatca tggaataacc tcaaagataa 1980
gcgccccatt gctaaacaat aacaaaaaat tattaagccg actgaaaaat tcaagacata 2040
ttagcactgg tgccatattg aataacacaa tcgcgactat aagcacaaat ccgaacttga 2100
attctaatgt gatgcagaac aataacaatc tgatgtcggg acacaatcac ctggacgagt 2160
tgagcagtat aaaacaggag ccaccgcatc aattgcagca gcagcaacca ccaatggatg 2220
tacaatcggt agattcgtat acctctgaca acccagacag caatgttatt gccaagtcgc 2280
ctgataagag gtcaagctta gtatccctat ctaaagtttc tccacattta ctttcatcca 2340
catcaagcaa cggtaacaca atatcgtgtc ccaatgttgc cacaaattcg caggaattgg 2400
aaccaaacaa tgatatttca acgaagaaat ctctttccaa ttctactttg agacattcct 2460
ctgctaatag aaattctaat tatggtgaca acaaaaggcc tcttagaaca acagtgtcaa 2520
agatatttga ttcaaacct aatggagetc ctttacggag atactctggg gtaccggatc 2580
acgttaatct agaagattac atcgaacagc cgcataatta tccaacaatg caaaatagtg 2640
tgaaaaagga tgaattttat aacagcagga acaataagtt tccccatggt ttaaactttt 2700
atggtgataa caatgttatt gaagaggaaa ataatggtga ctcgtctaat gtaaatcgac 2760
cgcaacacac taaccttcag catgagttta ttccagaaga taacgaaagt gatgaaaacg 2820
atattcactc catgttttat tataatcata agaacgattt agaaacaaaa ccgctaatat 2880
ccgattatgg tgaagatgaa gacgtagatg attatgatcg cccaaatgct actttcaaca 2940
gttactatgg ctcagcatcc aacacgcacg aacttccatt acatggaagg atgccttcaa 3000
gatcaaataa tgattactac gattttatgg ttggcaacaa tactggcaat aacaaccaat 3060
tgaatgaata taccccctta agaatgaaac gtggtcaaag acacctatca agaacaaaca 3120
atagcataat gaatggtagc atccatatga atggtaacga tgacgttacc cattccaata 3180
tcaataataa cgatattgtt ggttactcac cgcacaactt ttactcaagg aagtccccat 3240
ttgtgaaagt aaagaatttt ctttatcttg catttgttat atcatcacta ttgatgacag 3300
gattcattct gggattttta ttggccacta ataaagaact acaagatgta gacgtggtag 3360
tgatggataa tgtgatttca agttcggacg agttgatctt cgacatcaca gtaagtgctt 3420
ttaatccagg attcttcagt ataagcgttt cccaagtcga tttggacatt tttgcaaaaa 3480
gttcttacct gaagtgcgat tctaatggtg actgtacagt aatggaacag gaacggaaaa 3540
ttttacaaat aacgacaaat ctttcgttag ttgaagagag tgctaataat gatattagtg 3600
gtgggaacat agagacggta ttactaggaa ccgctaaaaa actagagaca ccattaaagt 3660
tccagggcgg cgcatttaat aggaactacg atgtgtcagt ctcgagtgtc aagcttttaa 3720
gtcctgggtc tcgtgaagcc aagcacgaaa acgacgatga tgacgatgat gatggcgacg 3780
atggtgacga tgaaaacaat actaatgaaa gacaatacaa aagcaaacca aatgctagag 3840
atgacaaaga agatgatact aaaaaatgga agctactaat caagcatgat tacgaattga 3900
tagtccgtgg aagcatgaag tatgaggtgc cctttttcaa tacgcaaaaa tctacggcta 3960
                                                                  3998
ttcaaaagga ttccatggtc catcctggta agaagtga
```

```
<210> 228
<211> 1165
<212> PRT
<213> Saccharomyces cerevisiae
```

Ala Pro Val Ala Asn Asn Leu Leu Ser Asn Asn Ser Asn Val Val

20

30

Ala Pro Asn Pro Ser Ile Pro Ser Ala Ser Thr Ser Thr Ser Pro Leu 35 40 45

His Arg Glu Ile Val Asp Asp Ser Val Ala Thr Ala Asn Thr Thr Ser 50 55 60

Asn Val Val Gln His Asn Leu Pro Thr Ile Asp Asn Asn Leu Met Asp 65 70 75 80

Ser Asp Ala Thr Ser His Asn Gln Asp His Trp His Ser Asp Ile Asn 85 90 95

Arg Ala Gly Thr Ser Met Ser Thr Ser Asp Ile Pro Thr Asp Leu His
100 105 110

Leu Glu His Ile Gly Ser Val Ser Ser Thr Asn Asn Asn Ser Asn Asn 115 120 125

Ala Leu Ile Asn His Asn Pro Leu Ser Ser His Leu Ser Asn Pro Ser 130 135 140

Ser Ser Leu Arg Asn Lys Lys Ser Ser Leu Leu Val Ala Ser Asn Pro 145 150 155 160

Ala Phe Ala Ser Asp Val Glu Leu Ser Lys Lys Lys Pro Ala Val Ile 165 170 175

Ser Asn Asn Met Pro Thr Ser Asn Ile Ala Leu Tyr Gln Thr Ala Arg 180 185 190

Ser Ala Asn Ile His Gly Pro Ser Ser Thr Ser Ala Ser Lys Ala Phe 195 200 205

Arg Lys Ala Ser Ala Phe Ser Asn Asn Thr Ala Pro Ser Thr Ser Asn 210 215 220

Asn Ile Gly Ser Asn Thr Pro Pro Ala Pro Leu Leu Pro Leu Pro Ser 225 230 235 240

Leu Ser Gln Gln Asn Lys Pro Lys Ile Ile Glu Arg Pro Thr Met His
245 250 255

Val Thr Asn Ser Arg Glu Ile Leu Leu Gly Glu Asn Leu Leu Asp Asp 260 265 270

Thr Lys Ala Lys Asn Ala Pro Ala Asn Ser Thr Thr His Asp Asn Gly

280

285

Pro Val Ala Asn Asp Gly Leu Arg Ile Pro Asn His Ser Asn Ala Asp 290 295 300

Asp Asn Glu Asn Asn Lys Met Lys Lys Asn Lys Asn Ile Asn Ser 305 310 315 320

Gly Lys Asn Glu Arg Asn Asp Asp Thr Ser Lys Ile Cys Thr Thr Ser 325

Thr Lys Thr Ala Pro Ser Thr Ala Pro Leu Gly Ser Thr Asp Asn Thr 340 345 350

Gln Ala Leu Thr Ala Ser Val Ser Ser Ser Asn Ala Asp Asn His Asn 355 360 365

Asn Asn Lys Lys Lys Thr Ser Ser Asn Asn Asn Gly Asn Asn Ser Asn 370 375 380

Ser Ala Ser Asn Lys Thr Asn Ala Asp Ile Lys Asn Ser Asn Ala Asp 385 390 395 400

Leu Ser Ala Ser Thr Ser Asn Asn Asn Ala Ile Asn Asp Asp Ser His
405 410 415

Glu Ser Asn Ser Glu Lys Pro Thr Lys Ala Asp Phe Phe Ala Ala Arg
420 425 430

Leu Ala Thr Ala Val Gly Glu Asn Glu Ile Ser Asp Ser Glu Glu Thr
435 440 445

Phe Val Tyr Glu Ser Ala Ala Asn Ser Thr Lys Asn Leu Ile Phe Pro 450 455 460

Asp Ser Ser Ser Gln Gln Gln Gln Gln Gln Gln Gln Pro Pro Lys Gln 465 470 475 480

Gln Gln Gln Gln Asn His Gly Ile Thr Ser Lys Ile Ser Ala Pro 485 490 495

Leu Leu Asn Asn Asn Lys Lys Leu Leu Ser Arg Leu Lys Asn Ser Arg 500 505 510

His Ile Ser Thr Gly Ala Ile Leu Asn Asn Thr Ile Ala Thr Ile Ser 515 520 525

Thr Asn Pro Asn Leu Asn Ser Asn Val Met Gln Asn Asn Asn Leu

SEQL

530

535

540

Met Ser Gly His Asn His Leu Asp Glu Leu Ser Ser Ile Lys Gln Glu 550 555 Pro Pro His Gln Leu Gln Gln Gln Pro Pro Met Asp Val Gln Ser 565 570 Val Asp Ser Tyr Thr Ser Asp Asn Pro Asp Ser Asn Val Ile Ala Lys 585 Ser Pro Asp Lys Arg Ser Ser Leu Val Ser Leu Ser Lys Val Ser Pro 595 600 His Leu Leu Ser Ser Thr Ser Ser Asn Gly Asn Thr Ile Ser Cys Pro 610 615 620 Asn Val Ala Thr Asn Ser Gln Glu Leu Glu Pro Asn Asn Asp Ile Ser 625 630 635 Thr Lys Lys Ser Leu Ser Asn Ser Thr Leu Arg His Ser Ser Ala Asn

645 650 655

Arg Asn Ser Asn Tyr Gly Asp Asn Lys Arg Pro Leu Arg Thr Thr Val 660 665 670

Ser Lys Ile Phe Asp Ser Asn Pro Asn Gly Ala Pro Leu Arg Arg Tyr 675 680 685

Ser Gly Val Pro Asp His Val Asn Leu Glu Asp Tyr Ile Glu Gln Pro 690 695 700

His Asn Tyr Pro Thr Met Gln Asn Ser Val Lys Lys Asp Glu Phe Tyr 705 710 715 720

Asn Ser Arg Asn Asn Lys Phe Pro His Gly Leu Asn Phe Tyr Gly Asp
725 730 735

Asn Asn Val Ile Glu Glu Glu Asn Asn Gly Asp Ser Ser Asn Val Asn 740 745 750

Arg Pro Gln His Thr Asn Leu Gln His Glu Phe Ile Pro Glu Asp Asn 755 760 765

Glu Ser Asp Glu Asn Asp Ile His Ser Met Phe Tyr Tyr Asn His Lys 770 775 780

Asn Asp Leu Glu Thr Lys Pro Leu Ile Ser Asp Tyr Gly Glu Asp Glu

790

795

800

Asp Val Asp Asp Tyr Asp Arg Pro Asn Ala Thr Phe Asn Ser Tyr Tyr 805 815

Gly Ser Ala Ser Asn Thr His Glu Leu Pro Leu His Gly Arg Met Pro 820 825 830

Ser Arg Ser Asn Asn Asp Tyr Tyr Asp Phe Met Val Gly Asn Asn Thr 835 840 845

Gly Asn Asn Asn Gln Leu Asn Glu Tyr Thr Pro Leu Arg Met Lys Arg 850 855 860

Gly Gln Arg His Leu Ser Arg Thr Asn Asn Ser Ile Met Asn Gly Ser 865 87,0 875 880

Ile His Met Asn Gly Asn Asp Asp Val Thr His Ser Asn Ile Asn Asn 885 890 895

Asn Asp Ile Val Gly Tyr Ser Pro His Asn Phe Tyr Ser Arg Lys Ser 900 905 910

Pro Phe Val Lys Val Lys Asn Phe Leu Tyr Leu Ala Phe Val Ile Ser 915 920 925

Ser Leu Leu Met Thr Gly Phe Ile Leu Gly Phe Leu Leu Ala Thr Asn 930 935 940

Lys Glu Leu Gln Asp Val Asp Val Val Val Met Asp Asn Val Ile Ser 945 950 955 960

Ser Ser Asp Glu Leu Ile Phe Asp Ile Thr Val Ser Ala Phe Asp Pro 965 970 975

Gly Phe Phe Ser Ile Ser Val Ser Gln Val Asp Leu Asp Ile Phe Ala 980 985 990

Lys Ser Ser Tyr Leu Lys Cys Asp Ser Asn Gly Asp Cys Thr Val Met 995 1000 1005 -

Glu Gln Glu Arg Lys Ile Leu Gln Ile Thr Thr Asn Leu Ser Leu Val 1010 1015 1020

Glu Glu Ser Ala Asn Asn Asp Ile Ser Gly Gly Asn Ile Glu Thr Val 1025 1030 1035 1040

Leu Leu Gly Thr Ala Lys Lys Leu Glu Thr Pro Leu Lys Phe Gln Gly

SEQL

1045

1050

1055

Gly Ala Phe Asn Arg Asn Tyr Asp Val Ser Val Ser Ser Val Lys Leu 1060 1065 1070

Leu Ser Pro Gly Ser Arg Glu Ala Lys His Glu Asn Asp Asp Asp Asp 1075 1080 1085

Asp Asp Asp Gly Asp Asp Gly Asp Asp Glu Asn Asn Thr Asn Glu Arg 1090 1095 1100

Gln Tyr Lys Ser Lys Pro Asn Ala Arg Asp Asp Lys Glu Asp Asp Thr 1105 1110 1115 1120

Lys Lys Trp Lys Leu Leu Ile Lys His Asp Tyr Glu Leu Ile Val Arg 1125 1130 1135

Gly Ser Met Lys Tyr Glu -Va-1- Pro Phe Phe Asn Thr Gln Lys Ser Thr 1140 1145 1150

Ala Ile Gln Lys Asp Ser Met Val His Pro Gly Lys Lys 1155 1160 1165

<210> 229

<211> 1076

<212> DNA

<213> Saccharomyces cerevisiae

<400> 229

gacgactatt gatgccaggc aaattttgga tttactgctc ctcttttaag aagacaagtg 60 tgtgatatcg tagcggtagg aaccaatttt gcaatcgatt tacttacagc caagaaaatc 120 tattttcatg tttagcattg ccatttcttc tgtgtcacac gttgtgcttg ccaggaacta 180 taggagagac gtatacaagc atcaatgtta cgaatgtacg atcccgtttg catctgatgt 240 gtaaactcat gtggtgcact ggtgttgttt ccaagactgc actattaact gggaattttt 300 ttttttcttc tagtgaattt ttttttaaag cgacgcacag gaaaagtgaa aattatttaa 360 acqqacqqca aacatqaaaa aaaaaattac caaccatatt tctatttcct ttccctttac 420 ctattctctt tttgaaatag ttcattttct ctctctgaaa cgacaataaa ccaaactcta 480 gcctccaata gtcactaaag atgaagtaca ttcaaaccga acaacaaatt gaaatcccag 540 aaggtgttac tgtcagcatt aagtccagaa tcgtcaaggt tgtcggtcca agaggtactt 600 tgaccaagaa cttgaagcat attgatgtta ccttcaccaa ggtcaacaac caattgatca 660 aggttgctgt tcacaacggt gacagaaagc acgttgccgc tttgagaacc gttaaatctt 720 tggttgacaa catgatcact ggtgtcacca agggttacaa gtacaagatg agatacgtct 780 acgcgcattt cccaatcaac gtcaacattg ttgaaaagga tggtgctaaa ttcattgaag 840 tcagaaactt tttgggtgac aagaagatca gaaacgtccc agttagagat ggtgttacta 900 tcgaattctc tactaacgta aaggacgaaa tcgtcttatc tggtaactct gttgaagacg 960 tttcccaaaa tgccgctgac ttgcaacaaa tctgtcgtgt tagaaacaag gatatccgta 1020

agtttttgga tggtatetac gtttcccaca agggtttcat tgtcgaagac atgtaa

1076

<210> 230

<211> 191

<212> PRT

<213> Saccharomyces cerevisiae

<400> 230

Met Lys Tyr Ile Gln Thr Glu Gln Gln Ile Glu Ile Pro Glu Gly Val 1 5 10 15

Thr Val Ser Ile Lys Ser Arg Ile Val Lys Val Val Gly Pro Arg Gly 20 25 30

Thr Leu Thr Lys Asn Leu Lys His Ile Asp Val Thr Phe Thr Lys Val 35 40 45

Asn Asn Gln Leu Ile Lys Val Ala Val His Asn Gly Asp Arg Lys His 50 55 60

Val Ala Ala Leu Arg Thr Val Lys Ser Leu Val Asp Asn Met Ile Thr 65 70 75 80

Gly Val Thr Lys Gly Tyr Lys Tyr Lys Met Arg Tyr Val Tyr Ala His.
85 90 95

Phe Pro Ile Asn Val Asn Ile Val Glu Lys Asp Gly Ala Lys Phe Ile 100 105 110

Glu Val Arg Asn Phe Leu Gly Asp Lys Lys Ile Arg Asn Val Pro Val 115 120 125

Arg Asp Gly Val Thr Ile Glu Phe Ser Thr Asn Val Lys Asp Glu Ile 130 135 140

Val Leu Ser Gly Asn Ser Val Glu Asp Val Ser Gln Asn Ala Ala Asp 145 150 155 160

Leu Gln Gln Ile Cys Arg Val Arg Asn Lys Asp Ile Arg Lys Phe Leu 165 170 175

Asp Gly Ile Tyr Val Ser His Lys Gly Phe Ile Val Glu Asp Met 180 185 190

<210> 231

<211> 1373

1373

```
<212> DNA
<213> Saccharomyces cerevisiae
<400> 231
tcgatggata tcccatccaa gaacaggaat actgggtttt tgaagaccag aatggagatc 60
tctgaggaag aaaagatggt acgtacaata tcacggcttg acaatacgag tattgcaaac 120
agtaatggaa atggtaatga tgacacctct aatcagagaa cggaagcact ggggcgtaag 180
acgagtaatg gagggcgaat atgattacta agttaaataa atcagataca gtatttaaag 240
tictttcaaa aaaagataat gtcatatatt ttactatcta cgcagtgaaa gagttccttc 300
taatgacaca ctattcactt cgggtaacgg atattgtgta ctgaaaaaata taaaaaattt 360
tatcccggaa atgcgatgag atgaaaatgc atgaagtagc gtatatattg attgcatgag 420
gttggacttg aaagggcata tatactcggt tttatcattg attcaagtgt tcccataaat 480
aataaaacag ttaaatcgaa atgctaagaa gacaagcccg tgaaaggaga gaatatctat 540
acagaaaagc gcaagaatta caagattctc aactgcaaca aaaacgtcaa ataattaaac 600
aagcgctagc tcaggggaag ccattgccaa aggaactagc agaagatgag agtttacaaa 660
aggatttcag atatgaccaa agtttaaagg agagcgaaga agcagatgat ctacaggttg 720
atgatgaata tgctgccaca agtggtataa tggatccaag aatcatcgtc acaacatctc 780
gtgacccaag cactcgtctc tcgcaatttg ccaaagaaat taaactgcta tttccaaatg 840
ctgtcaggct gaacagaggt aattatgtga tgccaaatct agtggatgct tgtaaaaaat 900
ccggtactac agatttggtg gtattacatg aacatagagg tgttccaact tctttgacca 960
atgatattat aaatgctggt aaccaaagcg aagtgaatcc acatctaata tttgataact 1080
ttactaccgc tttagggaaa agagtagtct gtattttaaa gcacttgttc aatgcggggc 1140
ccaaaaaaga ttccgaaaga gtaatcactt ttgcgaatag gggtgatttc attagcgtta 1200
gacagcatgt atatgtgaga acaagagagg gagtagagat tgccgaagtt ggtcctagat 1260
ttgagatgag gttgtttgaa ctgaggttgg gaactttaga aaataaggac gctgatgttg 1320
agtggcagtt gagaagattc ataaggactg ccaataaaaa agactatttg tga
<210> 232
<211> 290
<212> PRT
<213> Saccharomyces cerevisiae
<400> 232
Met Leu Arg Arg Gln Ala Arg Glu Arg Arg Glu Tyr Leu Tyr Arg Lys
                 5
                                    10
                                                       15
 1
Ala Gln Glu Leu Gln Asp Ser Gln Leu Gln Gln Lys Arg Gln Ile Ile
            20
                                                    30
Lys Gln Ala Leu Ala Gln Gly Lys Pro Leu Pro Lys Glu Leu Ala Glu
                                                45
                            40
        35
```

55

Asp Glu Ser Leu Gln Lys Asp Phe Arg Tyr Asp Gln Ser Leu Lys Glu

60

Ser Glu Glu Ala Asp Asp Leu Gln Val Asp Asp Glu Tyr Ala Ala Thr
65 70 75 80

Ser Gly Ile Met Asp Pro Arg Ile Ile Val Thr Thr Ser Arg Asp Pro 85 90 95

Ser Thr Arg Leu Ser Gln Phe Ala Lys Glu Ile Lys Leu Leu Phe Pro 100 105 110

Asn Ala Val Arg Leu Asn Arg Gly Asn Tyr Val Met Pro Asn Leu Val 115 120 125

Asp Ala Cys Lys Lys Ser Gly Thr Thr Asp Leu Val Val Leu His Glu 130 135 140

His Arg Gly Val Pro Thr Ser Leu Thr Ile Ser His Phe Pro His Gly
145 150 155 160

Pro Thr Ala Gln Phe Ser Leu His Asn Val Val Met Arg His Asp Ile 165 170 175

Ile Asn Ala Gly Asn Gln Ser Glu Val Asn Pro His Leu Ile Phe Asp 180 185 190

Asn Phe Thr Thr Ala Leu Gly Lys Arg Val Val Cys Ile Leu Lys His
. 195 200 205

Leu Phe Asn Ala Gly Pro Lys Lys Asp Ser Glu Arg Val Ile Thr Phe 210 215 220

Ala Asn Arg Gly Asp Phe Ile Ser Val Arg Gln His Val Tyr Val Arg 225 230 235 240

Thr Arg Glu Gly Val Glu Ile Ala Glu Val Gly Pro Arg Phe Glu Met 245 250 255

Arg Leu Phe Glu Leu Arg Leu Gly Thr Leu Glu Asn Lys Asp Ala Asp 260 265 270

Val Glu Trp Gln Leu Arg Arg Phe Ile Arg Thr Ala Asn Lys Lys Asp 275 280 285

Tyr Leu 290

<210> 233

<211> 1418

```
<212> DNA
<213> Saccharomyces cerevisiae
<400> 233
aaacacctac ttatagacac gaccaaactt tccacaacct ttcatcagag agaaatgttg 60
atcaagttga atgcgtgaaa gtagcaattc gaaacaacaa ctacctgtca ttctgcatag 120
tagtagttac gaaaggcaca gaaaataaca aaaaaaaaa aaaagtcaat tttctacggt 180
ctccatccgt acctctttaa atccgtacat tattgttttg cttaatttca atatttcgga 240
aaaagcgagc gccctggtaa aatgtggttc aagcctgcga gcctttgctt ggtaactcac 300
caaatgcaat tcagtcacgt tccacacagt ttgggtttcc agcctggctt tagggaagaa 360
tgggctcact aggcgttcat aatacgcgga gggggaaata ccaaatgcta ttgattatgg 420
ttaaaatatg tgttatttga ctttgtatat acaaacagaa gagaaaccaa cacactaaag 480
gtgagttgga attacaagtc gccaagacct tcatcgatct agaaagctcc tctccagaac 600
taaaggctga cttgagacca ttgcaaatca aatctatcag agaagtatgt taaaagttat 660
ataatttgga agcagcaaca ttgtgatttc ttctaaaggg gttctttgca gtaatttttt 720
caaaaaagag tgattttgag cagtatctgt atgaaatttt catgtgttcg agaaaaatag 780
taattccgag agctgtcaat accatgaacg ttgcgatgag cctttgaact ataaaggcct 840
ccttggtcag taccaatatc gatgaataaa atagaagcac gcgaaaaaga ccttacccca 900
aggagaagaa tcacaaaccc ttttttgtta tgaatgaacc aattcagtta ctaactttat 960
ttcaacgctg cttgattctt attgtttaga ttgatgtcac cggtggtaag aaagcactag 1020
tectttttgt eccagtteca getttgtetg cataceataa ggtecaaace aaattgacee 1080
gtgaattgga aaagaaattc cctgaccgtc atgttatttt cttggctgaa agaagaatct 1140
tgccaaaacc atctagaaca tctagacaag tccaaaagag accaagatcc agaactttga 1200
ctgctgttca cgacaaggtt ttggaagaca tggttttccc aactgaaatt gtcggtaaaa 1260
gagttagata tttggttggt ggtaacaaga tccaaaaggt tttgttagac tccaaggatg 1320
ttcaacaaat cgactacaag ttggaatctt tccaagctgt ctacaacaag ttgactggca 1380
aacaaattgt ttttgaaatt ccaagccaga ccaactaa
                                                                1418
<210> 234
<211> 190
<212> PRT
<213> Saccharomyces cerevisiae
<400> 234
Met Ser Ser Val Gln Ser Lys Ile Leu Ser Gln Ala Pro Ser Glu Leu
 1
                                    10
Glu Leu Gln Val Ala Lys Thr Phe Ile Asp Leu Glu Ser Ser Ser Pro
                                25
Glu Leu Lys Ala Asp Leu Arg Pro Leu Gln Ile Lys Ser Ile Arg Glu
                            40
Ile Asp Val Thr Gly Gly Lys Lys Ala Leu Val Leu Phe Val Pro Val
```

60

55

```
Pro Ala Leu Ser Ala Tyr His Lys Val Gln Thr Lys Leu Thr Arg Glu
65
                   70
Leu Glu Lys Lys Phe Pro Asp Arg His Val Ile Phe Leu Ala Glu Arg
                                  90
               85
Arg Ile Leu Pro Lys Pro Ser Arg Thr Ser Arg Gln Val Gln Lys Arg
           100
                              105
Pro Arg Ser Arg Thr Leu Thr Ala Val His Asp Lys Val Leu Glu Asp
                  120 125
```

Met Val Phe Pro Thr Glu Ile Val Gly Lys Arg Val Arg Tyr Leu Val 140

Gly Gly Asn Lys Ile Gln Lys Val Leu Leu Asp Ser Lys Asp Val Gln 150 155

Gln Ile Asp Tyr Lys Leu Glu Ser Phe Gln Ala Val Tyr Asn Lys Leu 170 165

Thr Gly Lys Gln Ile Val Phe Glu Ile Pro Ser Gln Thr Asn 190 180 185

135

<210> 235 <211> 1333 <212> DNA <213> Saccharomyces cerevisiae

<400> 235

130

ttcatcacca atatagacta atgcgttttg gaacgccaaa ccgcagtgac aaatagcaaa 60 tatgtagctg tcatatcggc atataataac agttttctac caaatgctgt cctacattca 120 gagatettae ateettaeat etaaagtaaa acetagaeat ttaettegag ttataetttt 180 tttttattta tctattttt ctcttgcgga catttaacac ctgaattccg cctaacgcca 240 ggactgatcc tgccagggaa gggagctttg tctagtgcca ataggccgga ccagtaggaa 300 ggttacagca gctggcccgc agagtgattg ggtcacagga aatagcgcaa ccttctcttt 360 tgcccgggaa aggcggttca atctaccttc gaagggctag tacatgagcg cgaaggaggc 420 agataatagc accattaagt ggtccaaatg catcttgaaa tctaatcctt aatagaggaa 480 aacaacaatt atcagtaaaa atgggtatgt tataaccata attcctaatg gtgaataaaa 540 tcaggaccaa taaagaaaag ctaatttgat ttttattgtc aatgaaattt cataatcgtc 600 atgaatgcat aaacagacac acctagcaac tgtataatct gcgcctaaaa agggcgtata 660 cacaaaacta aacgatgcgc aataaaagtt cagcagtcag caatgaaacc gagatatgca 720 gcaacagagt atcatatgca tggaggatcc tttctgtttt tctgataata tgctctgaaa 780 aagctccaaa cagcacagta gcctatttgt gaagctcaaa aaaggcttct atttccctcg 840 ctatcttcag attgtgcagt gatattcttt gaggaaggaa acgtagaggg gataagttgg 900

```
ataactgtta tttctttca atatgctaga ttttgcttac caccttactg atttttcta 960 ataataaact tttttactaa cattagtacg atgtctcatc tatttcttct atttagttaa 1020 cgttccaaag accagaaaga cctactgtaa gggtaagacc tgtcgtaagc acactcaaca 1080 caaggttact caatacaaag ctggtaaggc ttccttgttt gcccaaggta agagacgtta 1140 tgaccgtaaa caatctggtt tcggtggtca aaccaagcct gtttccaca agaaagctaa 1200 gactaccaag aaggttgttt tgagattgga atgtgtcaaa tgtaagacca gagcccaatt 1260 gaccttgaag agatgcaagc acttcgaat gggtggtgaa aagaagcaaa agggtcaagc 1320 tttgcaattc tga
```

```
<210> 236
```

<211> 116

<212> PRT

<213> Saccharomyces cerevisiae

<400> 236

Met Val Arg Cys Leu Ile Tyr Phe Phe Tyr Leu Val Asn Val Pro Lys

1 5 10 15

Thr Arg Lys Thr Tyr Cys Lys Gly Lys Thr Cys Arg Lys His Thr Gln 20 25 30

His Lys Val Thr Gln Tyr Lys Ala Gly Lys Ala Ser Leu Phe Ala Gln
35 40 45

Gly Lys Arg Arg Tyr Asp Arg Lys Gln Ser Gly Phe Gly Gln Thr
50 55 60

Lys Pro Val Phe His Lys Lys Ala Lys Thr Thr Lys Lys Val Val Leu 65 70 75 80

Arg Leu Glu Cys Val Lys Cys Lys Thr Arg Ala Gln Leu Thr Leu Lys
85 90 95

Arg Cys Lys His Phe Glu Leu Gly Gly Glu Lys Lys Gln Lys Gly Gln
100 105 110

Ala Leu Gln Phe 115

<210> 237

<211> 1223

<212> DNA

<213> Saccharomyces cerevisiae

<400> 237

```
ggtccacgtc agttccaeac aataacattt acgtagtgtt cacgcgaagc agttacatct 60
caactaacat aattgctggt gagcctacaa cactgcatgc gtaaacgtca acgggattac 120
gttagtattt ttggccgccg gtaaattctc ttgttttttt ttcttgattt cacttctttt 180
catgttcctt tggaataatc taattcctca tgattaaatg agactgtttt ttgtttccgt 240
aacatccata cctttcctgt ataatattct tgctgtaaag tttgtttttt ttatgaaaaa 300
aacattttct tttcttgaga tgaggcgccg cgagcctttc tcccatgggc agtggtaaat 360
tttccaaatc aatgcagctc tttgaaatac aacagcattt ttcatacatt ttaagcaatt 420
tctagtttgt agatattgtt agattagttt ttgaacattg ttttgataac tgaaaataaa 480
acagcaaaca aactacaaaa atggtcgctt taatctctaa gaaaagaaag ctagtcgctg 540
acggtgtctt ctacgctgaa ttgaacgaat tcttcaccag agaattagct gaagaaggtt 600
actocggtgt tgaagtocgt gtoactocaa ccaagaccga agttatoatc agagctacca 660
gaactcaaga tgttttgggt gaaaacggta gaagaatcaa cgaattaact ttgttggttc 720
aaaagagatt caagtacgct ccaggtacta ttgtcttata tgctgaaaga gttcaagacc 780
gtggtttgtc cgctgtcgct caagctgaat ctatgaaatt caaattgttg aacggtttgg 840
ctatcagaag agctgcttac ggtgtcgtca gatacgttat ggaatctggt gctaagggtt 900
gtgaagttgt tgtttccggt aaactaagag ctgccagagc taaggctatg aaatttgctg 960
acggtttctt gattcactct ggtcaaccag tcaacgactt cattgacact gctactagac 1020
acgtcttgat gagacaaggt gttttgggta tcaaggttaa gattatgaga gacccagcta 1080
agagcagaac tggtccaaag gctttgccag atgctgtcac catcattgaa ccaaaagaag 1140
aagaaccaat tettgeteca tetgteaagg actacagace agetgaagaa actgaagete 1200
aagctgaacc agttgaagct tag
                                                                  1223
```

<210> 238

<211> 240

<212> PRT

<213> Saccharomyces cerevisiae

<400> 238

Met Val Ala Leu Ile Ser Lys Lys Arg Lys Leu Val Ala Asp Gly Val

1 5 10 15

Phe Tyr Ala Glu Leu Asn Glu Phe Phe Thr Arg Glu Leu Ala Glu Glu 20 25 30

Gly Tyr Ser Gly Val Glu Val Arg Val Thr Pro Thr Lys Thr Glu Val
35 40 45

Ile Ile Arg Ala Thr Arg Thr Gln Asp Val Leu Gly Glu Asn Gly Arg
50 55 60

Arg Ile Asn Glu Leu Thr Leu Leu Val Gln Lys Arg Phe Lys Tyr Ala 65 70 75 80

Pro Gly Thr Ile Val Leu Tyr Ala Glu Arg Val Gln Asp Arg Gly Leu 85 90 95

Ser Ala Val Ala Gln Ala Glu Ser Met Lys Phe Lys Leu Leu Asn Gly

SEQL

100

105

110

Leu Ala Ile Arg Arg Ala Ala Tyr Gly Val Val Arg Tyr Val Met Glu 115 120 125

Ser Gly Ala Lys Gly Cys Glu Val Val Val Ser Gly Lys Leu Arg Ala 130 135 140

Ala Arg Ala Lys Ala Met Lys Phe Ala Asp Gly Phe Leu Ile His Ser 145 150 155 160

Gly Gln Pro Val Asn Asp Phe Ile Asp Thr Ala Thr Arg His Val Leu
165 170 175

Met Arg Gln Gly Val Leu Gly Ile Lys Val Lys Ile Met Arg Asp Pro 180 185 190

Ala Lys Ser Arg Thr Gly Pro Lys Ala Leu Pro Asp Ala Val Thr Ile 195 200 205

Ile Glu Pro Lys Glu Glu Glu Pro Ile Leu Ala Pro Ser Val Lys Asp 210 215 220

Tyr Arg Pro Ala Glu Glu Thr Glu Ala Gln Ala Glu Pro Val Glu Ala
225 230 235 240

<210> 239

<211> 2168

<212> DNA

<213> Saccharomyces cerevisiae

<400> 239

ctttgataaa ttaatacggt aagataccgt gtgaactatt ataataactg ccacgcttat 60 agcatgtacg ctatacattt acgtgctgag ctcctaggaa agctcatgag cagccactgt 120 atcgtggagc ataactacaa caaagaatac acagcgtcac atagagggtt tttgagaggga 180 gaagttgaaa taggacttga tcttggggga gaggggattt gaaagcaccc attcaggagt 240 atgtgtctgt aattgaagtg ttagcgcgcg attcacctgt aataagagtg atgatttgat 300 agcgccattc tacatcata ggcaaatgtt gaaaaactgt acgcgcgaac taaaattttt 360 ttttacatcc cactaaatga aaattttaaa tcgatgccca ttccaaatat gcttattcga 420 aggacggctc tgacaagggc atatgcgtta agattgattg ttcaatattc ataaaacagg 480 atctttcaag ggacgataaa atggatgagc aagttattt tacaacaaat acctcaggaa 540 caatagcttc tgtacactca tttgaacaga taaattttaa tccttttat tgctcaagca caaaaagcat 660 gaaatagctg tgttcaagta ggaaataaat acctttttat tgctcaagca caaaaagcat 660

```
taatcaatgt ctacaatctg tcaggttctt tcaaaagaga atctgttgaa cagcgcttac 720
cattacctga aatcctaaaa tgtctggaag tagttgaaaa tgatggtgtg cagtatgata 780
gaattcaagg tgtcaatcat aatttaccag acttcaatct tccgtacctt ttacttggct 840
ccaccgaatc gggtaaattg tacatatggg agttaaattc agggatttta ttgaacgtga 900
agcctatggc tcattaccaa agtatcacca agattaagtc cattttaaac ggcaagtata 960
ttattacttc tggtaacgat tcgagagtta ttatatggca aactgttgac ttggtatcag 1020
cgtccaatga tgatcctaag cctttatgta tccttcacga tcatactcta cccgtgacag 1080
atttccaagt ttcttctagt caaggaaaat ttttatcatg tactgatacg aaactcttca 1140
cagtatetea agatgetace attagatget atgatttgag tttaatagge ageaaaaaga 1200
agcagaaggc aaacgaaaat gacgttagta ttggtaagac cccagtattg cttgcgacat 1260
ttacaactcc ttattctatc aaatccattg tactggatcc tgctgacaga gcatgctata 1320
ttggtactgc ggaaggttgt tittcattga atttatttta taaactaaag ggtaatgcta 1380
tegttaatet getacagtee geeggagtaa acacagttea aaaaggtagg gtttttteee 1440
tagtgcaacg taactcacta actggcggcg aaaatgaaga tttggatgca ctatatgcaa 1500
tgggccaact tgtctgtgag aatgtcctaa attcaaatgt gtcatgccta gaaatatcaa 1560
tggatggtac attattattg atcggtgata cggaggggaa agtttctatt gcggaaattt 1620
actcaaaaca aatcattaga actatccaaa ctttaactac atcacaggat tcagttggag 1680
aagtgaccaa tctcttaacc aacccttaca gactcgaacg tggaaattta ctttttgaag 1740
gagaatccaa aggcaaacaa cctagtaata ataatggtca caattttatg aagataccaa 1800
acttacaaag agttatcttt gatggtaaaa acaaaggcca tttacacgat atttggtatc 1860
agataggaga accagaagca gagacagatc ctaacctcgc attaccactt aacgacttta 1920
atgcctattt ggagcaggtc aaaacgcaag aatcgatatt ttcacatatc ggtaaggtgt 1980
caagcaatgt aaaagtgatt gacaataaaa tcgacgccac ttcatcttta gacagcaatg 2040
ccgctaaaga tgaggaaatt acagaactta agaccaacat agaagcatta actcatgcct 2100
acaaggagtt acgtgacatg cacgaaaagc tgtacgagga acaccaacag atgcttgaca 2160
                                                                  2168
agcaataa
```

```
<210> 240
```

<211> 555

<212> PRT

<213> Saccharomyces cerevisiae

### <400> 240

Met Asp Glu Gln Val Ile Phe Thr Thr Asn Thr Ser Gly Thr Ile Ala 1 5 10 15

Ser Val His Ser Phe Glu Gln Ile Asn Leu Arg Gln Cys Ser Thr Gln 20 25 30

Ser Arg Asn Ser Cys Val Gln Val Gly Asn Lys Tyr Leu Phe Ile Ala 35 40 45

Gln Ala Gln Lys Ala Leu Ile Asn Val Tyr Asn Leu Ser Gly Ser Phe
50 55 60

Lys Arg Glu Ser Val Glu Gln Arg Leu Pro Leu Pro Glu Ile Leu Lys 65 70 75 80

Cys	s Leu	. Glu	ı Val	. Val 85		Asn	Asp	Gly	Val 90		Tyr	Asp	Arg	Ile 95	Glr
Gly	√ Val	. Asn	His 100		Leu	Pro	Asp	Phe 105	Asn	Leu	Pro	Tyr	Leu 110	Leu	Let
Gly	Ser	Thr 115		Ser	Gly	Lys	Leu 120	Tyr	Ile	Trp	Glu	Leu 125	Asn	Ser	Gl
Ile	Leu		Asn	Val	Lys		Met	Ala	His	Tyr			Ile	Thr	Lys
	130					135		-			140				
Ile 145	Lys	Ser	Ile	Leu	Asn 150	Gly	Lys	Tyr	Ile	Ile 155	Thr	Ser	Gly	Asn	Asp 160
Ser	Arg	Val	Ile	Ile 165	Trp	Gln	Thr	Val	Asp 170	Leu	Val	Ser	Ala	Ser 175	Asr
Asp	Asp	Pro	Lys 180	Pro	Leu	Cys	Ile	Leu 185	His	Asp	His	Thr	Leu 190	Pro	Val
Thr	Asp	Phe 195	Gln	Val	Ser	Ser	Ser 200	Gln	Gly	Lys	Phe	Leu 205	Ser	Cys	Thr
Asp	Thr 210	Lys	Leu	Phe	Thr	Val 215	Ser	Gln	Asp	Ala	Thr 220	Ile	Arg	Cys	Туг
Asp 225	Leu	Ser	Leu	Ile	Gly 230	Ser	Lys	Lys	Lys	Gln 235	Lys	Ala	Asn	Glu	Asn 240
Asp	Val	Ser	Ile	Gly 245	Lys	Thr	Pro	Val	Leu 250	Leu	Ala	Thr	Phe	Thr 255	Thr
Pro	Tyr	Ser	Ile 260	Lys	Ser	Ile	Val	Leu 265	Asp	Pro	Ala	Asp	Arg 270	Ala	Cys
Tyr	Ile	Gly 275	Thr	Ala	Glu	Gly	Cys 280	Phe 	Ser	Leu	Asn	Leu 285	Phe	Tyr	Lys
Leu	Lys 290	Gly	Asn	Ala	Ile	Val 295	Asn	Leu	Leu	Gln	Ser 300	Ala	Gly	Val	Asn
	Val	Gln	Lys	Gly		Val	Phe	Ser	Leu		Gln	Arg	Asn	Ser	
305					310					315					320
Thr	Gly	Gly	Glu	Asn 325	Glu	Asp	Leu	Asp	Ala 330	Leu	Tyr	Ala	Met	Gly 335	Gln

Leu Val Cys Glu Asn Val Leu Asn Ser Asn Val Ser Cys Leu Glu Ile 340 345 350

Ser Met Asp Gly Thr Leu Leu Leu Ile Gly Asp Thr Glu Gly Lys Val 355 360 365

Ser Ile Ala Glu Ile Tyr Ser Lys Gln Ile Ile Arg Thr Ile Gln Thr 370 375 380

Leu Thr Thr Ser Gln Asp Ser Val Gly Glu Val Thr Asn Leu Leu Thr

Asn Pro Tyr Arg Leu Glu Arg Gly Asn Leu Leu Phe Glu Gly Glu Ser 405 410 415

Lys Gly Lys Gln Pro Ser Asn Asn Gly His Asn Phe Met Lys Ile 420 425 430

Pro Asn Leu Gln Arg Val Ile Phe Asp Gly Lys Asn Lys Gly His Leu 435 440 445

His Asp Ile Trp Tyr Gln Ile Gly Glu Pro Glu Ala Glu Thr Asp Pro 450 455 460

Asn Leu Ala Leu Pro Leu Asn Asp Phe Asn Ala Tyr Leu Glu Gln Val 465 470 475 480

Lys Thr Gln Glu Ser Ile Phe Ser His Ile Gly Lys Val Ser Ser Asn 485 490 495

Val Lys Val Ile Asp Asn Lys Ile Asp Ala Thr Ser Ser Leu Asp Ser 500 505 510

Asn Ala Ala Lys Asp Glu Glu Ile Thr Glu Leu Lys Thr Asn Ile Glu 515 520 525

Ala Leu Thr His Ala Tyr Lys Glu Leu Arg Asp Met His Glu Lys Leu 530 535 540

Tyr Glu Glu His Gln Gln Met Leu Asp Lys Gln 545 550 555

<210> 241

<211> 1115

<212> DNA

# <213> Saccharomyces cerevisiae

```
<400> 241
aatgcgctcc cgtacgtcag tggctgttgc tgaaacgaga caatttctca attcgtttgt 60
ttgtgtactg tatttgttat ctttactata tatatgttgt taagtttctt ttaccaatta 120
gtgctcactt ctctcgtctt ttattaggtg tgtgtgttgt gcgtaatttt cgtttcgctg 180
attactttat atagtgtagt ttgttcttga atgtaataaa gacttctgtt ttattttgtt 240
ttgttattta gaaacagtct atctggttta acttaaacga gtgagcttaa gataatctga 300
ctacaagaaa accaagcttc tattactttg tttctttctc ttttttcttt tttgaataaa 360
gaattttcct ttaaggagta acttaagcat ttagctgcac attaaacact ttttttttta 420
cttctaactc acacactttt ggaagaacat ttattttttc gaccttcttt cccaaatacc 480
cagogottta taattgaaat atgaagttot ottotgttac tgotattact ctagocacog 540
ttgccaccgt tgccactgct aagaagggtg aacatgattt cactaccact ttaactttgt 600
categgaegg tagtttaact actaceact etacteatac caeteacaag tatggtaagt 660
tcaacaagac ttccaagtcc aagaccccaa accacatgg tactcacaag tacggtaagt 720
tcaacaagac ctccaagtct aagaccccaa accataccgg tactcacaag tatggtaagt 780
tcaacaagac ttccaagtcc aagactccaa accataccgg tactcacaag tacggtaagt 840
tcaacaagac ctccaaatcc aagactccaa accacactgg tactcacaag tacggtaagt 900
tcaacaagac Ctccaagtct aagaccccaa accataccgg tactcacaag tatggtaagt 960
tcaacaaaac caaacatgac actaccactt atggtcctgg tgaaaaggcc cgtaagaaca 1020
atgccgccc tggtccatct aatttcaact ccataaaatt gtttggtgtt accgctggta 1080
gtgctgccgt agccggtgcc ttattactat tataa
                                                                  1115
```

<210> 242

<211> 204

<212> PRT

<213> Saccharomyces cerevisiae

<400> 242

Met Lys Phe Ser Ser Val Thr Ala Ile Thr Leu Ala Thr Val Ala Thr 1 5 10 15

Val Ala Thr Ala Lys Lys Gly Glu His Asp Phe Thr Thr Leu Thr
20 25 30

Leu Ser Ser Asp Gly Ser Leu Thr Thr Thr Thr Ser Thr His Thr Thr 35 40 45

His Lys Tyr Gly Lys Phe Asn Lys Thr Ser Lys Ser Lys Thr Pro Asn 50 55 60

His Thr Gly Thr His Lys Tyr Gly Lys Phe Asn Lys Thr Ser Lys Ser
65 70 75 80

Lys Thr Pro Asn His Thr Gly Thr His Lys Tyr Gly Lys Phe Asn Lys 85 90 95

02-07-1999

Thr Ser Lys Ser Lys Thr Pro Asn His Thr Gly Thr His Lys Tyr Gly
100 105 110

Lys Phe Asn Lys Thr Ser Lys Ser Lys Thr Pro Asn His Thr Gly Thr
115 120 125

His Lys Tyr Gly Lys Phe Asn Lys Thr Ser Lys Ser Lys Thr Pro Asn 130 135 140

His Thr Gly Thr His Lys Tyr Gly Lys Phe Asn Lys Thr Lys His Asp 145 150 155 160

Thr Thr Thr Tyr Gly Pro Gly Glu Lys Ala Arg Lys Asn Asn Ala Ala 165 170 175

Pro Gly Pro Ser Asn Phe Asn Ser Ile Lys Leu Phe Gly Val Thr Ala 180 185 190

Gly Ser Ala Ala Val Ala Gly Ala Leu Leu Leu Leu 195 200

<210> 243

<211> 1115

<212> DNA

<213> Saccharomyces cerevisiae

# <400> 243

ggttatacac atatatatat ttttcatttt taatgtctta gcttttgtat cttagatgaa 60 gttttagttc tgtatatcac gatcaagata tcatacaatc ataaattcaa ttattcttct 120 gtttcccctc ttgaggcatc aaacgagtgt ttgactgata cacaccaaca tactaaggca 180 actititing ctgcccaaag ctgtggcacg tatgaaactg ctititinget gcataaaaca 240 accatgtgga gtttttactg tattcgcatt tcgccccgct agcattcttc gttcatgcta 300 aaaatgaggc gtgggctaat attcagtatt aataattccg gcacccgcac agcccatacc 360 ggaaaagggg ctggctgttg ggcttggcaa aaaactcaat ctgagcagtc atttataaag 420 aaagacttta atttgtcttg ctaaacactt gtaagccttc caaatataga tcacttaaga 480 caatctaaca agtgtccaaa atgtctgcaa acgaattcta ctcaagtggc caacaaggtc 540 aatataacca gcaaaacaac caagaaagaa ctggtgctcc aaacaacggt caatatggtg 600 ccgacaatgg taaccccaac ggtgaacgtg gtttattttc cactattgta ggtggcagtg 660 ccggtgcgta cgctggatct aaggtgtcga acaaccattc taagttgagt ggtgtgctgg 720 gcgccatagg tggtgcattc cttgccaaca agatatctga tgagcgtaaa gagcataagc 780 aacaagagca atacggcaac tcaaacttcg gaggtgctcc tcaaggtgga cacaacaacc 840 atcaccgtca gacaataaca acaataacgg tggatttggc ggtccaggcg gccctggcgg 900 tcaaggtttc ggaagacaag gcccacaagg atttggaggt cctggtccac aagagtttgg 960 tggtccaggt ggccaaggat tcggtggtcc aaatcctcaa gaattcggcg gccaggtggc 1020 caaggattcg gtggtccaaa ccctcaggaa ttcgggggcc aaggtcgtca aggattcaat 1080 ggcggttcac gttggtgaat ggctcaacag agtga 1115

<2	1	U.	24	lZ

<211> 204

<212> PRT

<213> Saccharomyces cerevisiae

<400> 244

Met Ser Ala Asn Glu Phe Tyr Ser Ser Gly Gln Gln Gly Gln Tyr Asn

1 5 10 15

Gln Gln Asn Asn Gln Glu Arg Thr Gly Ala Pro Asn Asn Gly Gln Tyr
20 25 30

Gly Ala Asp Asn Gly Asn Pro Asn Gly Glu Arg Gly Leu Phe Ser Thr
35 40 45

Ile Val Gly Gly Ser Ala Gly Ala Tyr Ala Gly Ser Lys Val Ser Asn 50 55 60

Asn His Ser Lys Leu Ser Gly Val Leu Gly Ala Ile Gly Gly Ala Phe 65 70 75 80

Leu Ala Asn Lys Ile Ser Asp Glu Arg Lys Glu His Lys Gln Gln Glu 85 90 95

Gln Tyr Gly Asn Ser Asn Phe Gly Gly Ala Pro Gln Gly Gly His Asn 100 105 110

Asn His His Arg Gln Thr Ile Thr Thr Ile Thr Val Asp Leu Ala Val 115 120 125

Gln Ala Ala Leu Ala Val Lys Val Ser Glu Asp Lys Ala His Lys Asp 130 135 140

Leu Glu Val Leu Val His Lys Ser Leu Val Val Gln Val Ala Lys Asp 145 150 155 160

Ser Val Val Gln Ile Leu Lys Asn Ser Ala Ala Arg Trp Pro Arg Ile 165 170 175

Arg Trp Ser Lys Pro Ser Gly Ile Arg Gly Pro Arg Ser Ser Arg Ile 180 185 190

Gln Trp Arg Phe Thr Leu Val Asn Gly Ser Thr Glu 195 200

```
<210> 245
<211> 1313
<212> DNA
<213> Saccharomyces cerevisiae
<400> 245
tcataacggg ttcttttcaa aaaaccgtaa aaatttgagg tcacaccaac taaatacaaa 60
ttgtttcatc acggtgacta tatcaagaac ttcgtaagga aacatttaga aaactcaata 120
tagtaaagtt tcatcagcaa tcttatctga gtaatattat ctacgatcta aatataggat 180
gatetgeega tttaggaate gtaetgtaga ttgetettgg egacagatat agtgaaatac 240
<del>cttttacaaa gtggatacag gttgcctatc actaccgcca tttcactagc aagtagagta 300</del>
ttgagaaaac ggtaaacttt gaaagttgca gatgcagaat atatatctgg ttttgtagtt 360
ctatccgcta aacgggacga tcgcatttta gccgccgaca gtgttaatat aagtaatgaa 420
cttgggttaa tttgattacg cgtcacagct actaataaaa taagaccgag agttttaatc 480
agctagtgca taccaaaaca atgagtaacc aacacagccc tcagccattt tgtttggaca 540
ccaaattggt gaaactatta gaagagctcc aggagggaaa gcaattcaac aataaaaaca 600
tattcccgga aaaagcatta tatttgaagc tcgctcttga ttattctttc ttcagaaaga 660
atttactaga gttttgcgtc caccttgaca agataaaagg agtcattaga ccaaactatg 720
acactatata tattttgtgc ctgttggagg tggatctcct caatctggta tttaccgaca 780
atatattgga aatatgtttg cccaggtttg tttcaaggga ggacttgagg gtttttaata 840
atacttttta cacatatcac gataaccgcc tacgtattct ccaagaagac ttttctcaat 900
tgttcaaaaa aatcaaaact aaggcttctg tactatgttt tacagttgag gaaatttttc 960
tgacaaacca agaaatttta cctcaaaact caacagtggc agaactgcaa aagagcacta 1020
ataaagtaca gacaaatggg ccgcaacggc acgatttcat agtcactcta gaaataaaac 1080
tgaacaaaac acaaatcact ttcctcattg gagctaaagg aacgagaatt gaaagcttga 1140
gggaaaaatc aggcgccagc ataaaaataa tacctattag tgataaaatg actgcacatg 1200
aaaggaacca ccctgaatct gttcaacaaa caatactaat ttcgggtgac ttatactcaa 1260
ttgcattagc cgtcaccagt atagagtctg cattaattac tttggattta tag
<210> 246
<211> 270
<212> PRT
<213> Saccharomyces cerevisiae
<400> 246
Met Ser Asn Gln His Ser Pro Gln Pro Phe Cys Leu Asp Thr Lys Leu
  1
                  5
                                      10
Val Lys Leu Leu Glu Glu Leu Gln Glu Gly Lys Gln Phe Asn Asn Lys
             20
                                                      30
Asn Ile Phe Pro Glu Lys Ala Leu Tyr Leu Lys Leu Ala Leu Asp Tyr
         35
                             40
                                                  45
Ser Phe Phe Arg Lys Asn Leu Leu Glu Phe Cys Val His Leu Asp Lys
    50
                         55
                                              60
```

1	Ù
	•

Ile	Lys	Gly	Val	Ile	Arg	Pro	Asn	Tyr	Asp	Thr	Ile	Tyr	Ile	Leu	Cys
65					70					75					80

Leu Leu Glu Val Asp Leu Leu Asn Leu Val Phe Thr Asp Asn Ile Leu 85 90 95

Glu Ile Cys Leu Pro Arg Phe Val Ser Arg Glu Asp Leu Arg Val Phe
100 105 110

Asn Asn Thr Phe Tyr Thr Tyr His Asp Asn Arg Leu Arg Ile Leu Gln
115 120 125

Glu Asp Phe Ser Gln Leu Phe Lys Lys Ile Lys Thr Lys Ala Ser Val 130 135 140

Leu Cys Phe Thr Val Glu Glu Ile Phe Leu Thr Asn Gln Glu Ile Leu 145 150 155 160

Pro Gln Asn Ser Thr Val Ala Glu Leu Gln Lys Ser Thr Asn Lys Val 165 170 175

Gln Thr Asn Gly Pro Gln Arg His Asp Phe Ile Val Thr Leu Glu Ile 180 185 190

Lys Leu Asn Lys Thr Gln Ile Thr Phe Leu Ile Gly Ala Lys Gly Thr 195 200 205

Arg Ile Glu Ser Leu Arg Glu Lys Ser Gly Ala Ser Ile Lys Ile Ile 210 215 220

Pro Ile Ser Asp Lys Met Thr Ala His Glu Arg Asn His Pro Glu Ser 225 230 235 240

Val Gln Gln Thr Ile Leu Ile Ser Gly Asp Leu Tyr Ser Ile Ala Leu 245 250 255

Ala Val Thr Ser Ile Glu Ser Ala Leu Ile Thr Leu Asp Leu 260 265 270

<210> 247

<211> 1766

<212> DNA

<213> Saccharomyces cerevisiae

<400> 247

```
agttttttt ctcgagaaat tgtgaacaaa agaaagcaaa gacacagaag atgataagag 60
agagaaacaa cgaagaaaga acaacaatgt tggggttcac ccgagagata ttgacatact 120
gaccttagaa aaggcattac tgaggctact gactaaagcg cgttacataa atgcatagta 180
tatttcttgt tgtatacgca gcggccaact agtggcagca agaatgtaat gaacgattca 240
tctgcaggtt tggaggccgc aactagatca aaacgtaaat agcgggtgaa gtgttctgga 300
cgttagaagt aacgtccgca gatcgaagct aaacacgaga ttagatttcg ggtaacggaa 360
ttgtgataat taagaaagac cagactatgt gaaaaggcca cgtaaatgat agagcacaca 420
ttagcaacta taatagacta gttttcgcat cgctggaagt tctcgatatt gaatatcact 480
tccaagaacg caaacttaga atggtccgga ttcttcccat aattttgagc gccctatctt 540
cgaaattagt ggcgagtaca atattgcatt catccataca ctcagtgcca tctggaggcg 600
aaatcatatc tgcagaagat cttaaagaac ttgaaatttc agggaattcg atctgcgttg 660
ataatcgttg ctatcctaag atatttgaac caagacacga ttggcagccc atactgccag 720
qtcaaqaact ccccggtggt ttggacatta gaataaacat ggacacaggt ttaaaagagg 780
caaaactaaa tgatgagaag aatgtcggtg ataatggtag ccatgagtta attgtatctt 840
cagaagacat gaaagcatcg cctggtgact atgaattttc cagtgatttc aaagaaatga 900
gaaacatcat agattctaac ccgactttat cttcacagga cattgccaga ttggaggata 960
gttttgatag aataatggaa tttgcgcatg attacaagca cggctacaaa attattaccc 1020
atgaattcgc cctcttggcc aaccttagtc tcaatgaaaa tttgccgtta acattgagag 1080
ageteagtac tagagteatt accagetget tgagaaacaa teeteetgta gtegagttea 1140
ttaatgaaag ttttccaaat tttaaaagca aaatcatggc cgctctgtca aatttgaatg 1200
attctaacca cagatcctct aatatcctaa taaaaagata cttgtccatt ttaaacgaat 1260
tacctgtcac atccgaagat cttcctatat actctacggt tgttttacaa aatgtatatg 1320
aaagaaacaa caaggacaaa cagttacaaa taaaagtcct ggagttgatc agcaaaattt 1380
tgaaggccga catgtacgaa aatgacgata caaatctaat tttgttcaaa agaaatgctg 1440
agaattggtc gtcaaatctg caagagtggg caaacgagtt ccaagagatg gtccagaaca 1500
aaagtataga tgaactacat acaagaacgt tttttgacac cctttacaac ttgaagaaaa 1560
ttttcaaaag tgacatcacg atcaacaaag ggtttttgaa ttggttagcg caacaatgta 1620
aagccaggca atctaacttg gacaatgggc tccaagagag agatactgaa caagactcat 1680
ttgataagaa acttatcgac agcagacact tgatctttgg caaccccatg gctcatagaa 1740
                                                                  1766
taaaaaattt cagagatgaa ctctga
```

```
<210> 248
```

<211> 421

<212> PRT

<213> Saccharomyces cerevisiae

<400> 248

Met Val Arg Ile Leu Pro Ile Ile Leu Ser Ala Leu Ser Ser Lys Leu
1 5 10 15

Val Ala Ser Thr Ile Leu His Ser Ser Ile His Ser Val Pro Ser Gly
20 25 30

Gly Glu Ile Ile Ser Ala Glu Asp Leu Lys Glu Leu Glu Ile Ser Gly
35 40 45

Asn Ser Ile Cys Val Asp Asn Arg Cys Tyr Pro Lys Ile Phe Glu Pro

55

60

Arg His Asp Trp Gln Pro Ile Leu Pro Gly Gln Glu Leu Pro Gly Gly 65 70 75 80

Leu Asp Ile Arg Ile Asn Met Asp Thr Gly Leu Lys Glu Ala Lys Leu 85 90 95

Asn Asp Glu Lys Asn Val Gly Asp Asn Gly Ser His Glu Leu Ile Val 100 105 110

Ser Ser Glu Asp Met Lys Ala Ser Pro Gly Asp Tyr Glu Phe Ser Ser 115 120 125

Asp Phe Lys Glu Met Arg Asn Ile Ile Asp Ser Asn Pro Thr Leu Ser 130 135 140

Ser Gln Asp Ile Ala Arg Leu Glu Asp Ser Phe Asp Arg Ile Met Glu 145 150 155 160

Phe Ala His Asp Tyr Lys His Gly Tyr Lys Ile Ile Thr His Glu Phe 165 170 175

Ala Leu Leu Ala Asn Leu Ser Leu Asn Glu Asn Leu Pro Leu Thr Leu 180 185 190

Arg Glu Leu Ser Thr Arg Val Ile Thr Ser Cys Leu Arg Asn Asn Pro 195 200 205

Pro Val Val Glu Phe Ile Asn Glu Ser Phe Pro Asn Phe Lys Ser Lys 210 215 220

Ile Met Ala Ala Leu Ser Asn Leu Asn Asp Ser Asn His Arg Ser Ser 225 230 235 240

Asn Ile Leu Ile Lys Arg Tyr Leu Ser Ile Leu Asn Glu Leu Pro Val 245 250 255

Thr Ser Glu Asp Leu Pro Ile Tyr Ser Thr Val Val Leu Gln Asn Val
260 265 270

Tyr Glu Arg Asn Asn Lys Asp Lys Gln Leu Gln Ile Lys Val Leu Glu 275 280 285

Leu Ile Ser Lys Ile Leu Lys Ala Asp Met Tyr Glu Asn Asp Asp Thr 290 295 300

Asn Leu Ile Leu Phe Lys Arg Asn Ala Glu Asn Trp Ser Ser Asn Leu

320

Gln Glu Trp Ala Asn Glu Phe Gln Glu Met Val Gln Asn Lys Ser Ile 325 330 335

Asp Glu Leu His Thr Arg Thr Phe Phe Asp Thr Leu Tyr Asn Leu Lys 340 345 350

Lys Ile Phe Lys Ser Asp Ile Thr Ile Asn Lys Gly Phe Leu Asn Trp 355 360 365

Leu Ala Gln Gln Cys Lys Ala Arg Gln Ser Asn Leu Asp Asn Gly Leu
370 375 380

Gln Glu Arg Asp Thr Glu Gln Asp Ser Phe Asp Lys Lys Leu Ile Asp 385 390 395 400

Ser Arg His Leu Ile Phe Gly Asn Pro Met Ala His Arg Ile Lys Asn 405 410 415

Phe Arg Asp Glu Leu 420

<210> 249

305

<211> 821

<212> DNA

<213> Saccharomyces cerevisiae

# <400> 249

taagtacatg attittgtti gcattgatat gacttgttti atgactaaca tatttaattt 60 ttatttgtta accgtagggg tittatgaag tgctgacgaa tcctgtttat tggaagcata 120 tittactgtt tgcggtttgc tatgccctga titttgtcac tattgctggt ctctittatg 180 tcacacttgt accgctitta gtgacatggg ccatactgtt attagggcct cttggtgtga 240 tactggttca tattcaatgg attitacaaa cgaatgtcti gactgcctit gittgtagaa 300 cactggtcct gacccatatt acgaatcaga tatttgatat atcittggtg titgcaagacc 360 aagatgaatt tctaaacgag gtgaaggtat tgcctaaacc acaaaagcca catagaaaaa 420 tcgatgaacc tgatgcggtg agaaatitca acacaataaa gggaagtcgg attittaaga 480 tcccagatt actaticaga atgittita aagictccaa tittactica ctaacattac 540 tgcgctaat tcctattgta ggaccaatci tggcaaatca actaatggcc ccaaaaagaa 600 ccttaccta titgcagagg tactittac taaagggati cagtaagaaa caggccaaag 660 attitcagta cgagcattac gcaagtitca tatgiticgg tatgictgg gcagctaaat 780 ggtgtacttc gctactaag ggtgaaagaa agaaggaatg a

<210> 250

SEQL

Arg Thr Phe Thr Tyr Leu Gln Arg Tyr Phe Leu Leu Lys Gly Phe Ser
35 40 45

Lys Lys Gln Ala Lys Asp Phe Gln Tyr Glu His Tyr Ala Ser Phe Ile 50 55 60

Cys Phe Gly Met Ser Ala Gly Leu Leu Glu Leu Ile Pro Phe Phe Thr
65 70 75 80

Ile Val Thr Ile Ser Ser Asn Thr Val Gly Ala Ala Lys Trp Cys Thr 85 90 95

Ser Leu Leu Lys Gly Glu Arg Lys Lys Glu 100 105

<210> 251 <211> 1256 <212> DNA <213> Saccharomyces cerevisiae

<400> 251

acatttecca aaaaagacat ttetgtecaa aagtagaagg caagaaaace etggaggaat 60 cataggeaaa gaaagaaaag aagaagttea tetttaaaac tacettteaa geetttatte 120 gtteetegta aaggacacae gaaaaaaata aacagtacet tgeagaagga gtgeagagtt 180 aggtegeagg gaateettga aageeaagag ttttttee gtaatgatet eecaaaageaa 240 eeateaacat tgtggtgeaa agtttagtgt aagatgttet aetgaaetat ettaataget 300 gageateatg tgagtaaacg agtaageaag aaaacaacaa agtaatgtte aactttegta 360 actaeggaaa ataatata agtagttaae gaaattegaa eaatgagage teteacatat 420 eatettett teeagtttag eeattateag eacaataata caaaacaac tegtacacte 480 getteaacta taacaaaaaa atggettaca teaagatege tttattaget getategetg 540 etttggette tgeecaaact eaggaagaaa ttgaegaat gaaegttatt ttgaatgaeg 600 ttaagteea ettgeagaa tatattagtt tggetgaaga ttetteatet ggattteet 660 taageagtet geeatetggt gttttagaea teggtttage tttggettee geeactgatg 720 acteetaca taetttgtae tetgaggttg actttgetge tgttageaag atgttgaeca 780 tggtteeatg gtattettee aggettetae eagaattgga ateettgtta ggaaetteta 840

ccaccgctgc ctcttctact gaagettett ctgctgctac ttcttccgct gttgcttcct 900 ccagtgaaac tacttettet gccgtcgctt cctccagtga agctacttet tcttgccgtcg 960 cttcttccag tgaagettet tcttctgctg ctacttette tgctgtcgct tcttccagtg 1020 aggetacete ttctcaget gcttcctcta ccaaggetge ctcttccact aaggettett 1080 cctctgctgt ttcttcaget gttgcttett ccaccaaage ctccgccatt tctcaaatca 1140 gtgatggtca agttcaagec actagcactg tttccgaaca aactgaaaac ggtgctgcca 1200 aggetgtcat cggtatgggt gctggtgtca tggccgctgc cgccatgtta ttataa 1256

<210> 252

<211> 251

<212> PRT

<213> Saccharomyces cerevisiae

<400> 252

Met Ala Tyr Ile Lys Ile Ala Leu Leu Ala Ala Ile Ala Ala Leu Ala 1 5 10 15

Ser Ala Gln Thr Gln Glu Glu Ile Asp Glu Leu Asn Val Ile Leu Asn 20 25 30

Asp Val Lys Ser Asn Leu Gln Glu Tyr Ile Ser Leu Ala Glu Asp Ser
35 40 45

Ser Ser Gly Phe Ser Leu Ser Ser Leu Pro Ser Gly Val Leu Asp Ile 50 55 60

Gly Leu Ala Leu Ala Ser Ala Thr Asp Asp Ser Tyr Thr Thr Leu Tyr
65 70 75 80

Ser Glu Val Asp Phe Ala Ala Val Ser Lys Met Leu Thr Met Val Pro 85 90 95

Trp Tyr Ser Ser Arg Leu Leu Pro Glu Leu Glu Ser Leu Leu Gly Thr
100 105 110

Ser Thr Thr Ala Ala Ser Ser Thr Glu Ala Ser Ser Ala Ala Thr Ser 115 120 125

Ser Ala Val Ala Ser Ser Glu Thr Thr Ser Ser Ala Val Ala Ser 130 135 140

Ser Ser Glu Ala Thr Ser Ser Ala Val Ala Ser Ser Ser Glu Ala Ser 145 150 155 160

Ser Ser Ala Ala Thr Ser Ser Ala Val Ala Ser Ser Ser Glu Ala Thr 165 170 175



Ser Ser Thr Val Ala Ser Ser Thr Lys Ala Ala Ser Ser Thr Lys Ala 180 185 190

Ser Ser Ser Ala Val Ser Ser Ala Val Ala Ser Ser Thr Lys Ala Ser 195 200 205

Ala Ile Ser Gln Ile Ser Asp Gly Gln Val Gln Ala Thr Ser Thr Val 210 215 220

Ser Glu Gln Thr Glu Asn Gly Ala Ala Lys Ala Val Ile Gly Met Gly 225 230 235 240

Ala Gly Val Met Ala Ala Ala Ala Met Leu Leu 245 250

<210> 253

<211> 2693

<212> DNA

<213> Saccharomyces cerevisiae

#### <400> 253

cattgatete gageaeaget getetttet eteaatgaat taegttatat gttaateaca 60 caagcatcag tttttcatcg caaaagaaaa tatttagagt tcttgcaatt cagatgtacc 120 tcaattaata actcaatcac ctcctatgtt cttgctggta gtactgcttt tgtcttaatt 180 actgctgaat cagccttcta gaagaccgtt ctgtttcagc cgctcgcccc ttttcaaagc 240 ttgcgcggct gagttttatg aggggcggct tttttgtgaa tggcaatcta ccattattag 300 tagcaatatg atttgcagat agatacatat atatccttct gggttcatgt ctcgttacca 360 tccacactaa tgcataggac cagaaagaaa ggacatcgaa tccaacacgt attaaaataa 420 ggactcctca ttaaaaaggc ttttaatatt caacttgcta tagatcacgc acccttttgg 480 caagtcaacc ttaaattatt atgatttctg tttgcccaca aaatgacttg caaaaatgct 540 acagaagcct cacattcgat gttccaggac aacaattcga agagagaaat gaacaaaacc 600 ttaaaaaacg ggccaaaaag aaaggcagtt tccaaccatc tgttgccttt gacacagtgc 660 cttccaccgc tggttattct tctatagacg acagcaggga aggattcaaa ggtgtacctg 720 ttcccaacta ttacacgatg gaagagtgct atgacgatga aacagactct ttttcgccaa 780 atttgcaata ttatttgaga gatacattcc aatcatcacc ttttctgaat actagaaaag 840 agaacaaatc tgaatccagt agttttccaa tgagatcctc aaagttgttg gaaaagaatt 900 ctgacatcaa aaaatatttc ttggtatcca agaatggaaa aatagtgagg agagactatc 960 caagcacgcc agtaattgtc aacgaaacgt tgatgataaa caggtttgaa aagaactgga 1020 taaagttatg gcgccaaaga aaactacaaa taaatgaaag gctgaatgac aaaaaaaaat 1080 ggtttactta cccagaactt atcttctctg aagagcgtat taaaccgtta tatagaggag 1140 atgatagtgc accatgtaca aaagaacaaa aaagaaagca taaaatactt caacaaaagg 1200 tcggatatcc caataaccct aagacaatag tttgtcacat taacggaaaa aaacatacgt 1260 gggttgccct agactggaca gtctacaagt ttgcacgaaa tcttgatcac attgttgtca 1320 taactacact gccaaaaatg atttctaaca ggaaaaaaac tgcaaaagat gatacagaat 1380 gggcaccggg atatcaaaaa gaagtaatag atcaaaaatt aaacgacatt tttgattata 1440 ttttacagct agtaaaagtg gtcaaaatat ccgtcaaaat tactttagaa ataattgtag 1500

```
gcaaaattaa aaaaagtetg gtagatgtea ttaatgteea tacteeagat ttettagtte 1560
ttgctacttt aaagcacgag cgaaatgaga atcttattac atataaatcc aaaaagctga 1620
cagatgtctt tcctgttagt tatccgattc ccacatttgt tgttccctcg aaacgaatgt 1680
attegttega aetgaateta caaagagaag taaatgaaca ttatgtetea aaaaateata 1740
tgaagcacga acacactgac gttgagagca tgagcagttc aatgttcaaa aaaaatacaa 1800
tatcagatat ttcttcacat atttccgtag attcgtacgc cgaagatttc aaaaggcaag 1860
gctacatcaa aaagcagttc aacacctcta atgattccat tccaagaaaa ttgaccggtc 1920
tcgcccagca ttcaagaagg aagatcacgg gtgatataga aaaattacaa gacgatgaga 1980
aagatagaga atgtactaag gaaaaacttt tgttgaagaa aattgatatc ataattagag 2040
agtcattgaa gtcttcttta gcgatagaga cgttgcctgg taaaaatgta tcgcagtcca 2100
gtcacggtga ccaaatttcc agctttaaga atgctttgat aggcaatggg tcgaaaaaca 2160
<del>caaagtttag aaaatettta ataeeatatt etteeteaga ggaacaaaat accacaacaa 222</del>0
ctattaaact cagtagctcg cctacgtccc aaatcaagtt tgcaacctct gtaaaacaca 2280
aagatggaag agccgccctt ggcaaagcca gaaatctgcc tgatataagg cacagtattt 2340
ccttcgacaa agaaaattcc tttgatccat ctgataaaag cagtagtgtt gataatagca 2400
ttcctttgag gaaagttaaa agtgccggtg cgttaagaaa agtcaaaact aatgactcct 2460
caagtagtgc agggtcaaag aaaagctcgt ctagttttag tactgtgaac accttcactg 2520
ggggtggagt tgggattttt aaggtgttta aaagtggaag ttcctctgga aataaatcat 2580
ccagtagaag gaatagtagc agtggcgatg tttttgaaag tgatgatcgt aacgacaaga 2640
aaaagaagaa gaaaaaaaag aagaaatcat tgttcttatt cggcaaaata tga
                                                                   2693
```

<210> 254

<211> 730

<212> PRT

<213> Saccharomyces cerevisiae

<400> 254

Met Ile Ser Val Cys Pro Gln Asn Asp Leu Gln Lys Cys Tyr Arg Ser 1 5 10 15

Leu Thr Phe Asp Val Pro Gly Gln Gln Phe Glu Glu Arg Asn Glu Gln 20 25 30

Asn Leu Lys Lys Arg Ala Lys Lys Lys Gly Ser Phe Gln Pro Ser Val 35 40 45

Ala Phe Asp Thr Val Pro Ser Thr Ala Gly Tyr Ser Ser Ile Asp Asp
50 55 60

Ser Arg Glu Gly Phe Lys Gly Val Pro Val Pro Asn Tyr Tyr Thr Met 65 70 75 80

Glu Glu Cys Tyr Asp Asp Glu Thr Asp Ser Phe Ser Pro Asn Leu Gln 85 90 95

Tyr Tyr Leu Arg Asp Thr Phe Gln Ser Ser Pro Phe Leu Asn Thr Arg 100 105 110

	пуs	GIU	115		261	Giu	Ser	120		Pile	PIO	Mec	125	Ser	ser	пÃ;
	Leu	Leu 130	Glu	Lys	Asn	Ser	Asp 135		Lys	Lys	Tyr	Phe 140	Leu	Val	Ser	Lys
	Asn 145	Gly	Lys	Ile	Val	Arg 150	Arg	Asp	Tyr	Pro	Ser 155	Thr	Pro	Val	Ile	Va]
_	Asn	Glu	Thr	Leu	Met 165		Asn	Arg	Phe	Glu <del>170</del>	_	Asn	Trp	Ile	Lys 175	
	Trp	Arg	Gln	Arg 180			Gln	Ile	Asn 185		٠	Leu		Asp 190	Lys	
	Lys	Trp	Phe 195	Thr	Tyr	Pro	Glu	Leu 200	Ile	Phe	Ser	Glu	Glu 205	Arg	Ile	Lys
	Pro	Leu 210	Tyr	Arg	Gly	Asp	Asp 215	Ser	Ala	Pro	Cys	Thr 220	Lys	Glu	Gln	Lys
	Arg 225	Lys	His	Lys	Ile	Leu 230	Gln	Gln	Lys	Val	Gly 235	Tyr	Pro	Asn	Asn	Pro 240
	Lys	Thr	Ile	Val	Cys 245	His	Ile	Asn	Gly	Lys 250	Lys	His	Thr	Trp	Val 255	Ala
	Leu	Asp	Trp	Thr 260	Val	Tyr	Lys	Phe	Ala 265	Arg	Asn	Leu	Asp	His 270	Ile	Val
	Val		Thr 275	Thr	Leu	Pro	Lys	Met 280	Ile	Ser	Asn	Arg	Lys 285	Lys	Thr	Ala
	Lys	Asp 290	Asp	Thr	Glu	Trp	Ala 295	Pro	Gly	Tyr	Gln	Lys 300	Glu	Val	Ile	Asp
	Gln 305	Lys	Leu	Asn	Asp	Ile 310	Phe	Asp	Tyr	Ile	Leu 315	Gln	Leu	Val	Lys	Val 320
	Val	Lys	Ile	Ser	Val 325	Lys	Ile	Thr	Leu	Glu 330	Ile	Ile	Val	Gly	Lys 335	Ile
	Lys	Lys	Ser	Leu 340	Val	Asp	Val	Ile	Asn 345	Val	His	Thr	Pro	Asp 350	Phe	Leu
	Val		Ala 355	Thr	Leu	Lys	His	Glu 360	Arg	Asn	Glu	Asn	Leu 365	Ile	Thr	Туг

Lys	Ser 370	Lys	Lys	Leu	Thr	Asp 375	Val	Phe	Pro	Val	Ser 380	Tyr	Pro	Ile	Pro
Thr 385	Phe	Val	Val	Pro	Ser 390	Lys	Arg	Met	Tyr	Ser 395	Phe	Glu	Leu	Asn	Leu 400
Gln	Arg	Glu	Val	Asn 405	Glu	His	Tyr	Val	Ser 410	Lys	Asn	His	Met	Lys 415	His
 Glu	His							Ser 425					Lys 430		Asn
Thr	Ile	Ser 435	Asp	Ile	Ser	Ser	His 440	Ile	Ser	Val	Asp	Ser 445	Tyr	Ala	Glu
Asp	Phe 450	Lys	Arg	Gln	Gly	туr 455	Ile	Lys	Lys	Gln	Phe 460	Asn	Thr	Ser	Asn
Asp 465	Ser	Ile	Pro	Arg	Lys 470	Leu	Thr	Gly	Leu	Ala 475	Gln	His	Ser	Arg	Arg 480
Lys	Ile	Thr	Gly	Asp 485	Ile	Glu	Lys	Leu	Gln 490	Asp	Asp	Glu	Lys	Asp 495	Arg
Glu	Cys	Thr	Lys 500	Glu	Lys	Leu	Leu	Leu 505	Lys	Lys	Ile	Asp	Ile 510	Ile	Ile
Arg	Glu	Ser 515	Leu	Lys	Ser	Ser	Leu 520	Ala	Ile	Glu	Thr	Leu 525		Gly	Lys
Asn	Val 530	Ser	Gln	Ser	Ser	His 535	Gly	Asp	Gln	Ile	Ser 540	Ser	Phe	Lys	Asn
Ala 545	Leu	Ile	Gly	Asn	Gly 550	Ser	Lys	Asn	Thr	Lys 555	Phe	Arg	Lys	Ser	Leu 560
										_				- 3 -	•

Ile Pro Tyr Ser Ser Ser Glu Glu Gln Asn Thr Thr Thr Thr Ile Lys 565 570 575

Leu Ser Ser Ser Pro Thr Ser Gln Ile Lys Phe Ala Thr Ser Val Lys 580 585 590

His Lys Asp Gly Arg Ala Ala Leu Gly Lys Ala Arg Asn Leu Pro Asp 595 600 605

Ile Arg His Ser Ile Ser Phe Asp Lys Glu Asn Ser Phe Asp Pro Ser 610 620

```
        Asp
        Lys
        Ser
        Ser
        Val
        Asp
        Asn
        Ser
        Ile
        Pro
        Leu
        Arg
        Lys
        Lys
        640

        Ser
        Ala
        Gly
        Ala
        Leu
        Arg
        Lys
        Val
        Lys
        Thr
        Asn
        Asp
        Ser
        Ser
        Ser
        Ser
        Ser
        655
        Ala
        Gly
        Ser
        Lys
        Ser
        Ser
        Ser
        Ser
        Phe
        Ser
        Thr
        Val
        Asn
        Thr
        Phe
        Fr
        Fr
```

Ser Gly Asn Lys Ser Ser Ser Arg Arg Asn Ser Ser Ser Gly Asp Val 690 695 700

Lys Lys Ser Leu Phe Leu Phe Gly Lys Ile 725 730

<210> 255 <211> 2270 <212> DNA <213> Saccharomyces cerevisiae

<400> 255

aattttcccc ccgtcataag ttcctataca cggctggctc tgatggcata atttcatgct 60 ggaacctaca aacccgcaag aaaataaaaa atttcgccaa atttaacgaa gacagcgtgg 120 ttaaaattgc ttgttcggac aatattctat gtctggcaac ttctgatgat actttcaaga 180 caaacgccgc aattgaccaa actattgaac taaacgcaag ttcaatatac ataatatttg 240 actatgagaa ctgatatctt cgtgaagatt cgtgtagtat gatagaacat tccagaaaaa 300 aaattcagat tcatcgctct ctcttcgctt ctcctccttt aaggaataaa gaaaaaatca 360 catacataga ttaagtaaat aggatctgct agaaaaatta tatatagatc aatcatctta 420 ttaaggtate ttgtttaage ceaaaagtet geteceaaat teeteactgt agetactaaa 480 acaacctata cgcaagaaag atgtcattga cagccgatga atacaaacaa caaggtaacg 540 ctgcatttac cgctaaggat tacgataaag cgatagagct cttcactaaa gctattgaag 600 tttctgaaac tccaaaccat gttttatatt ctaacaggtc cgcctgttat acttctttaa 660 agaaatttag tgacgcattg aatgatgcta atgaatgtgt caaaatcaat ccatcttggt 720 ctaagggtta taatagactc ggtgccgccc acttaggtct tggcgatctc gacgaagctg 780 aaagcaacta caaaaaagcc ttggagttgg atgccagtaa caaggccgcc aaagaaggat 840 tggatcaggt tcatcgtacc caacaggcaa gacaggcaca gcctgattta gggttgacac 900 agttgtttgc tgacccaaat ttaattgaaa atttaaagaa gaacccaaaa actagcgaaa 960 tgatgaagga ccctcaatta gtggctaaac tgattgggta caaacaaaat ccgcaagcta 1020 ttggccaaga tctgtttact gatccaagat taatgaccat catggctaca ttgatggggg 1080

```
ttgatttaaa catggatgat ataaaccaat caaactccat gccaaaggaa ccggaaacca 1140
qtaaaagcac tgaacaaaag aaagatgctg aaccacaaag cgattccact acgagcaagg 1200
aaaattcctc taaagcacca cagaaagaag aaagtaagga atccgagcca atggaagttg 1260
atgaagatga ctctaaaatt gaggccgaca aggaaaaggc cgaaggtaac aagttttaca 1320
aggcacgtca attcgatgaa gctatagagc actacaacaa ggcgtgggaa ctgcataaag 1380
atattaccta tttaaacaac cgtgctgctg ctgaatacga aaaaggcgaa tacgagacag 1440
ctatttctac cttgaatgat gctgttgagc aaggtagaga aatgagagcg gattacaagg 1500
tcatttccaa atcatttgcg cgtattggta atgcctatca caaattgggt gacttgaaga 1560
aaactataga atactaccaa aaatcattga ccgaacatcg tactgctgac attttgacca 1620
agttaaggaa tgctgaaaaa gaattgaaga aagctgaggc ggaggcgtat gttaaccctg 1680
aaaaggcgga ggaagcccgt cttgaaggta aggaatattt taccaagagt gattggccga 1740
atgetgttaa ggettacaet gaaatgatea aaagggeaee tgaagatget agaggatatt 1800
ctaatagagc tgctgcacta gcgaagttaa tgtctttccc tgaagctatc gcagattgta 1860
acaaagccat tgaaaaagat ccaaatttcg tgagagctta tatcagaaag gccaccgcac 1920
aaattgctgt taaagaatat gcttccgctt tggaaacact agatgcggcc agaaccaaag 1980
atgctgaagt gaataatggt tctagtgcaa gggaaattga tcaactgtac tacaaggcaa 2040
gccaacaaag attccaacct ggtaccagta acgaaacccc agaagaaacc tatcaaaggg 2100
ccatgaaaga tcctgaagtg gctgcgatca tgcaagatcc tgttatgcaa agtattttgc 2160
agcaggccca acagaatccc gctgctttac aagaacacat gaaaaatcca gaagtattca 2220
aaaagattea gaegttgate getgetggta teateeggae tggeegetaa
                                                                  2270
```

<210> 256

<211> 589

<212> PRT

<213> Saccharomyces cerevisiae

<400> 256

Met Ser Leu Thr Ala Asp Glu Tyr Lys Gln Gln Gly Asn Ala Ala Phe

1 5 10 15

Thr Ala Lys Asp Tyr Asp Lys Ala Ile Glu Leu Phe Thr Lys Ala Ile
20 25 30

Glu Val Ser Glu Thr Pro Asn His Val Leu Tyr Ser Asn Arg Ser Ala 35 40 45

Cys Tyr Thr Ser Leu Lys Lys Phe Ser Asp Ala Leu Asn Asp Ala Asn 50 55 60

Glu Cys Val Lys Ile Asn Pro Ser Trp Ser Lys Gly Tyr Asn Arg Leu 65 70 75 80

Gly Ala Ala His Leu Gly Leu Gly Asp Leu Asp Glu Ala Glu Ser Asn 85 90 95

Tyr Lys Lys Ala Leu Glu Leu Asp Ala Ser Asn Lys Ala Ala Lys Glu 100 105 110

G1;	y Let	Ası 11:	o Glr	ı Val	. His	arç	120		Gln	a Ala	Arg	Gln 125		Gln	Pro
Asj	p Let 130		, Leu	Thr	Gln	135		: Ala	Asp	Pro	Asn 140	Leu	Ile	Glu	Asn
Le:		Lys	s Asn	Pro	Lys 150		Ser	Glu	Met	Met 155		Asp	Pro	Gln	Leu 160
Va]	L Ala	Lys	Leu	lle		Tyr	Lys	Gln			Gln	Ala	Ile		
				103					<del>-170</del>					175	
Asp	) Leu	Phe	180		Pro	Arg	Leu	Met 185		Ile	Met	Ala	Thr 190	Leu	Met
Gly	v Val	Asp 195	Leu	Asn	Met	Asp	Asp 200	Ile	Asn	Gln	Ser	Asn 205	Ser	Met	Pro
Lys	Glu 210		Glu	Thr	Ser	Lys 215	Ser	Thr	Glu	Gln	Lys 220	Lys	Asp	Ala	Glu
Pro 225		Ser	Asp	Ser	Thr 230	Thr	Ser	Lys	Glu	Asn 235	Ser	Ser	Lys	Ala	Pro 240
Gln	Lys	Glu	Glu	Ser 245	Lys	Glu	Ser	Glu	Pro 250	Met	Glu	Val	Asp	Glu 255	Asp
Asp	Ser	Lys	Ile 260	Glu	Ala	Asp	Lys	Glu 265	Lys	Ala	Glu	Gly	Asn 270	Lys	Phe
Tyr	Lys	Ala 275	Arg	G1n	Phe	Asp	Glu 280	Ala	Ile	Glu	His	Tyr 285	Asn	Lys	Ala
Trp	Glu 290	Leu	His	Lys	Asp	Ile 295	Thr	Tyr	Leu	Asn	Asn 300	Arg	Ala	Ala	Ala
Glu 305	Tyr	Glu	Lys	Gly	Glu 310	Tyr	Glu	Thr	Ala	Ile 315	Ser	Thr	Leu	Asn	Asp 320
Ala	Val	Glu	Gln	Gly 325	Arg	Glu	Met	Arg	Ala 330	Asp	Tyr	Lys	Val	Ile 335	Ser
Lys	Ser	Phe	Ala	Arg	Ile	Gly	Asn	Ala	Tyr	His	Lys	Leu	Gly	Asp	Leu

360

Lys Lys Thr Ile Glu Tyr Tyr Gln Lys Ser Leu Thr Glu His Arg Thr

345

350

340

Ala Asp Ile Leu Thr Lys Leu Arg Asn Ala Glu Lys Glu Leu Lys Lys 370 375 380

Ala Glu Ala Glu Ala Tyr Val Asn Pro Glu Lys Ala Glu Glu Ala Arg 385 390 395 400

Leu Glu Gly Lys Glu Tyr Phe Thr Lys Ser Asp Trp Pro Asn Ala Val 405 410 415

Lys Ala Tyr Thr Glu Met Ile Lys Arg Ala Pro Glu Asp Ala Arg Gly
420 425 430

Tyr Ser Asn Arg Ala Ala Ala Leu Ala Lys Leu Met Ser Phe Pro Glu 435 440 445

Ala Ile Ala Asp Cys Asn Lys Ala Ile Glu Lys Asp Pro Asn Phe Val

Arg Ala Tyr Ile Arg Lys Ala Thr Ala Gln Ile Ala Val Lys Glu Tyr 465 470 475 480

Ala Ser Ala Leu Glu Thr Leu Asp Ala Ala Arg Thr Lys Asp Ala Glu 485 490 495

Val Asn Asn Gly Ser Ser Ala Arg Glu Ile Asp Gln Leu Tyr Tyr Lys
500 505 510

Ala Ser Gln Gln Arg Phe Gln Pro Gly Thr Ser Asn Glu Thr Pro Glu
515 520 525

Glu Thr Tyr Gln Arg Ala Met Lys Asp Pro Glu Val Ala Ala Ile Met 530 535 540

Gln Asp Pro Val Met Gln Ser Ile Leu Gln Gln Ala Gln Gln Asn Pro 545 550 555 560

Ala Ala Leu Gln Glu His Met Lys Asn Pro Glu Val Phe Lys Lys Ile 565 570 575

Gln Thr Leu Ile Ala Ala Gly Ile Ile Arg Thr Gly Arg 580 585

<210> 257

<211> 710

<212> DNA

```
<213> Saccharomyces cerevisiae
```

```
ctgcagaagt acagctgcct ttattcttg tggtcattta ttgcttttat tttcaagtca 60 gatatacaag aaaatcaaat cccatcgtca acgtcacgta taaacgatta atttacagta 120 ataccatact ctaccaacat tatttagtc cgacgttcag tcctgtaggt gttccaaatc 180 cttctggcat tgacttctgt gcagaaaccc ttcaaaatga gttccacttt acgtcagatc 240 gcataacaac cggtcatata ttttttctt ttgctaaacc ccctactgca agcactttta 300 agaaaaagga caataaatgc gtctttattg ctgtgtggaa gtgatttttg tctttcggac 360 aaaaaaagga tagggatgcg agagggctgt gaagtagtga tcaagcgggg cctatataag 420 aagggcgcac atcgtcccc ctaagaatag cgaagcgata ttacactgaa cactacaatg 480 tcaaatagta ctcaataaat atgactgtaa aaatatgtga ctgtgaaggc gaatgttgta 540 aggactcttg tcattgtggg agcacctgcc ttccaagctg ttctggcggt gaaaagtgca 600 aatgtgatca cagcaccgga agccctcaat gtaagagttg tggtgaaaaa tgcaaatgcg 660 aaaccacgtg cacttgtgaa aagagtaaat gcaattgtga aaaatgttag
```

<210> 258

<211> 69

<212> PRT

<213> Saccharomyces cerevisiae

<400> 258

Met Thr Val Lys Ile Cys Asp Cys Glu Gly Glu Cys Cys Lys Asp Ser 1 5 10 15

Cys His Cys Gly Ser Thr Cys Leu Pro Ser Cys Ser Gly Glu Lys
20 25 30

Cys Lys Cys Asp His Ser Thr Gly Ser Pro Gln Cys Lys Ser Cys Gly
35 40 45

Glu Lys Cys Lys Cys Glu Thr Thr Cys Thr Cys Glu Lys Ser Lys Cys
50 55 60

Asn Cys Glu Lys Cys 65

<210> 259

<211> 1474

<212> DNA

<213> Saccharomyces cerevisiae

<400> 259

aaacccatac acaatgaacc ttatcacacc caaacatatg atatggtatt aaaaaatgaa 60 aaaaattcat tattctttag cgtaattatt gaagaaaaaa cagtgcgcgc ggtaattttt 120

```
tgtcactcag taactagaga gaagccgaat gtactccccc ggctagctgg agaccatggc 180
tctgcctagg atttctctta tgctttcctt tcaccaatca ctttgttccg gcgaggcccg 240
cgaagetege tttetteag cetageaate atgttettge cagegtegta gaetactgta 300
tggcagttgc tgcacttgcc atgaatatcc tagtgaagcc tctatgcaat aatccagtta 360
ctgcgttaga atcctggtaa aatgtctaat cttattacat tacagcaacg tattagattt 420
tgattgaaaa ttagtccttg cgacttggta tatatcttat tttaagaaag ctgaaaggaa 480
gaaagatcat cacgaacaac atgtctgctc cacaagccaa gattttgtct caagctccaa 540
ctgaattgga attacaagtt gctcaagctt tcgttgaatt ggaaaattct tctccagaat 600
tgaaagctga gttgagacct ttgcaattca agtccatcag agaagtatgt tattaatttg 660
aatctaaact taagaataat ggagagtaac aaaggaaaaa agtgtgaacg ggacgatacc 720
agaatgtttc aatctagaaa agtataaaaag ataaggacta ggactcaaat gtatttggct 780
gactategee tgaacettga tgetaageaa ataceatate tteaagaaaa ageetaetee 840
agtgtttaag aagaagggaa cgatttacta gatcatgcta tacgcagtaa ggttctgata 900
gttaattaca atcggtccaa gttctaagcg gtgtcgtcca tgcatatatc atttacaagt 960
tactggcgtc aactettcaa atattcaaaa tatcacetaa teaaaettae taacatttte 1020
cttttttgtt ttccttcttt tatagatcga cgttgctggt ggtaagaagg ctttggccat 1080
ttttgttcca gtcccatctt tggctggttt ccacaaggtt caaactaagt tgacccgtga 1140
attggaaaag aaattccaag accgtcatgt catcttcttg gctgaaagaa gaatcttgcc 1200
aaagccatct agaacatcta gacaagtcca aaagagacca agatccagaa ccttgactgc 1260
tgttcatgac aagatcttgg aagacttagt cttcccaact gaaatcgttg gtaagagagt 1320
tagatatttg gttggtggta acaagatcca aaaggttttg ttagactcaa aggatgtcca 1380
acaaatcgac tacaaattgg aatctttcca agctgtttac aacaaattga ctggtaagca 1440
aattgttttc gaaattccaa gtgaaactca ttag
```

<210> 260

<211> 190

<212> PRT

<213> Saccharomyces cerevisiae

<400> 260

Met Ser Ala Pro Gln Ala Lys Ile Leu Ser Gln Ala Pro Thr Glu Leu

1 5 10 15

Glu Leu Gln Val Ala Gln Ala Phe Val Glu Leu Glu Asn Ser Ser Pro 20 25 30

Glu Leu Lys Ala Glu Leu Arg Pro Leu Gln Phe Lys Ser Ile Arg Glu
35 40 45

Ile Asp Val Ala Gly Gly Lys Lys Ala Leu Ala Ile Phe Val Pro Val
50 55 60

Pro Ser Leu Ala Gly Phe His Lys Val Gln Thr Lys Leu Thr Arg Glu 65 70 75 80

Leu Glu Lys Lys Phe Gln Asp Arg His Val Ile Phe Leu Ala Glu Arg 85 , 90 95

```
Arg Ile Leu Pro Lys Pro Ser Arg Thr Ser Arg Gln Val Gln Lys Arg 100 105 110

Pro Arg Ser Arg Thr Leu Thr Ala Val His Asp Lys Ile Leu Glu Asp 115 120 125

Leu Val Phe Pro Thr Glu Ile Val Gly Lys Arg Val Arg Tyr Leu Val 130 135 140

Gly Gly Asn Lys Ile Gln Lys Val Leu Leu Asp Ser Lys Asp Val Gln 145 150 155 160
```

Gln Ile Asp Tyr Lys Leu Glu Ser Phe Gln Ala Val Tyr Asn Lys Leu 165 170 175

Thr Gly Lys Gln Ile Val Phe Glu Ile Pro Ser Glu Thr His 180 185 190

```
<210> 261
<211> 803
<212> DNA
<213> Saccharomyces cerevisiae
```

<400> 261
acccatttta caaattttt ttgctattg agccatagta cccattaata ggtctcgtcc 60
attcccttgt tttttttta ttgtttcaat tacactacat aattaaaaat cacatcactt 120
tcactctcac cttagtcgtt ctttatcaac caaaaataaa aaaatgcttc aatccgttgt 180
ctttttcgct cttttaacct tcgcaagttc tgtgtcagcg atttattcaa acaatactgt 240
ttctacaact accactttag cgccagcta ctccttggtg ccccaagaga ctaccatatc 300
gtacgccgac gacaccacta ccttttttgt cacctcaacg gtctactcca cgagctggtt 360
cacctcaact tcagccaca ttaccaatgc ggcctcctcc tccttgtcca cctcttcggc 420
etctggatct gtaaccccag aatccacca tgaaattacc tccacctcga ctatcacgtc 480

cacctcaact tcagccacca ttaccaatgc ggcctcctc tccttgtcca cctcttcggc 420 ctctggatct gtaaccccag aatccacca tgaaattacc tccacctcga ctatcacgtc 480 cactttgctg ctaacccttc atgactccac tactttgtct ccatcatcta ctgcagcaag 540 tgtcagtgac gaagattcaa acaacaaaga tgcaaaggtc aagtcctttg aacaggcttc 600 aacttccaat ggttgcgtcc caatcacaaa gtttgtcact gtcaccaatg agcccgttac 660 ccagtacgtt acagtcaccc caaatacgac tacacaatac gttactgtca ccggtgcacc 720

ttctgttacc actacctctc caggtaacgt acaatggtac aacaccactt cgattactaa 780 ttcgaccagt tggtgaatta tga 803

<210> 262 <211> 100 <212> PRT

<213> Saccharomyces cerevisiae

```
<400> 262
```

Met Thr Pro Leu Leu Cys Leu His His Leu Leu Gln Gln Val Ser Val
1 5 10 15

Thr Lys Ile Gln Thr Thr Lys Met Gln Arg Ser Ser Pro Leu Asn Arg
20 25 30

Leu Gln Leu Pro Met Val Ala Ser Gln Ser Gln Ser Leu Ser Leu Ser 35 40 45

Pro Met Ser Pro Leu Pro Ser Thr Leu Gln Ser Pro Gln Ile Arg Leu

50 55 60

His Asn Thr Leu Leu Ser Pro Val His Leu Leu Pro Leu Pro Leu 65 70 75 80

Gln Val Thr Tyr Asn Gly Thr Thr Pro Leu Arg Leu Leu Ile Arg Pro 85 90 95

Val Gly Glu Leu 100

<210> 263

<211> 1255

<212> DNA

<213> Saccharomyces cerevisiae

## <400> 263

aactttcgaa atatatactc tgagttcttg ctccccgttt tgtacgatgt tcctaatgca 60 aacacacgtc ctttaaagat cccctgatag gtttcattaa aggcaacttc catacacgtt 120 gaagtgccaa ttttttccct acatccaagc attctgggtt tgtatgggtg ttacaccggt 180 ttttcttttt tatttccaga gaagtacaat tttaggcggg tttcaaaattt cccctgtgtg 240 cgagaaacgc tccggtacgc ctaggctcac tccggtcctt ctccccattt ctatcagcgc 300 gataggcata ctgtgggaga gcggcactag ggagaccggt gggaagcacc gtatctagta 360 acquacquic tittgcaaaa atatccatta attgcatgta acttagatta acactggtat 420 taagatttcg caattttggg ctggattatt aaggtcgagt agcaaagttt agcaagaaca 480 gtacgaacta agtagccaag atgttgatgc caaaggaaga cagaaacaag atccaccaat 540 acttattcca aggtatgttt tagaataact ttcagaaagc atgaagatac acggaaagtc 600 aagcgaggga agttatgcgt atacacagta gcggtagtgt gcattcacat acacgatgtt 660 tcaaacacac agatggatac catgcatatg aggttaaagg atttcttatg aatatattag 720 tggattacat agaagaaatt acaaggaacc gtgtgacgac attttcgaaa ggacagcaca 780 aggccatcga ccctggagac gtatgaaata tggtatacgt cctatatttg ggcaagaaaa 840 cqqtaaaact gttttacgcg gagatccaac ttttgatcct taccgagtac cacgaatcat 900 ctatatgate tttttactaa ettatettea attteaaegg agagatgtaa acateattet 960 ctcctatgat aatttctttt tttatacaga aggtgttgtt gtcgccaaga aggatttcaa 1020 ccaagccaag cacgaagaaa ttgacaccaa gaacttgtat gtcattaagg ctttacaatc 1080 cttgacttct aagggttacg tcaagactca attctcatgg caatactact actacacctt 1140 gactgaagaa ggtgttgaat acttgagaga atacttgaac ttgccagaac acattgttcc 1200 aggtacctac attcaagaaa gaaacccaac tcaaagacca caaagaagat attaa 1255

<210> 264

<211> 105

<212> PRT

<213> Saccharomyces cerevisiae

### <400> 264

Met Leu Met Pro Lys Glu Asp Arg Asn Lys Ile His Gln Tyr Leu Phe 1 5 10 15

Gln Glu Gly Val Val Val Ala Lys Lys Asp Phe Asn Gln Ala Lys His
20 25 30

Glu Glu Ile Asp Thr Lys Asn Leu Tyr Val Ile Lys Ala Leu Gln Ser 35 40 45

Leu Thr Ser Lys Gly Tyr Val Lys Thr Gln Phe Ser Trp Gln Tyr Tyr 50 55 60

Tyr Tyr Thr Leu Thr Glu Glu Gly Val Glu Tyr Leu Arg Glu Tyr Leu 65 70 75 80

Asn Leu Pro Glu His Ile Val Pro Gly Thr Tyr Ile Gln Glu Arg Asn 85 90 95.

Pro Thr Gln Arg Pro Gln Arg Arg Tyr 100 105

<210> 265

<211> 1432

<212> DNA

<213> Saccharomyces cerevisiae

## <400> 265

tatttacaag ctagataaaa aaaaaatcaa atagcaagct attctgtcat atcttaaggt 60 gctgattgtt tgtaggcacg aagtaaacaa gctcttcccc aaaacaataa atacggtttt 120 cgggactgtt cacccgtaca ttttacgttt ccgacggcgc gatgctatct ttgaattttt 180 atctctttca ggtaacccca tgaccagtag ggcggcttac taccgaagag aaaaatgtcc 240 gcggcctaga cagttacttc ccaggccagg gccaggccac acggacagag gcagattcca 300 agttgttccg catagtctgt ctagctcttt ctcaattttc cgccagattc tgtctaattt 360 ctttccgctc gagttggcaa cagtacgaag aagtaactct aatagataga tataaccgtt 420 tttgagggca ttatttttgc agaagataat agaagagaac cgtaacaaag gaatcaagca 480

```
aagaaaaagt atgtaataga atgtattgta tgcttgtgga atgaatgtca atgtagctat 540
tttatatggt gagctcaaat tgaatgaaca tatcgtggaa atttaaaata ctgaagaata 600
cccaataagt caatgcaacc tgtgaatgtt tttcctgaaa tacgccgaat actgaatacg 660
attattacca taaattgctt ctagaaggaa ggcggtgtac catttaatac tgatgatatg 720
gttaatatca tttgagaagt cttcacatga ggactataga actaccatcc aggaattata 780
gaggaaatta actgaatcag agatctattt tgaaacattc atttacatgt aattgtctgc 840
aataaagcaa tatttttgaa atatgcaagt ttactaacaa gaataaattc ttttttgatt 900
tttatcttta acagtggctc atttcaaaga ataccaagtc attggtcgtc gtttaccaac 960
tgaatccgtt ccagaaccaa agttgttcag aatgagaatt tttgcttcaa atgaagtcat 1020
cgccaagtet cgttactggt atttcttgca aaaattgcac aaggttaaga aggettetgg 1080
tgaaattgtt tccatcaacc aaatcaacga agctcaccca accaaggtca agaacttcgg 1140
tgtttgggtt agatacgatt ccagatctgg tactcacaac atgtacaagg aaatcagaga 1200
cgtatccaga gttgctgccg tcgaaacctt ataccaagac atggctgcta gacacagagc 1260
tagatttaga tetatteaca tettgaaggt tgetgaaatt gaaaagactg etgatgteaa 1320
gagacaatac gtcaagcaat tcttgaccaa ggatttaaaa ttcccattac ctcacagagt 1380
ccaaaaatct accaagactt tctcttacaa gagaccatca accttctact aa
```

```
<210> 266
```

<211> 174

<212> PRT

<213> Saccharomyces cerevisiae

<400> 266

Met Tyr Leu Ala His Phe Lys Glu Tyr Gln Val Ile Gly Arg Arg Leu
1 5 10 15

Pro Thr Glu Ser Val Pro Glu Pro Lys Leu Phe Arg Met Arg Ile Phe
20 25 30

Ala Ser Asn Glu Val Ile Ala Lys Ser Arg Tyr Trp Tyr Phe Leu Gln
35 40 45

Lys Leu His Lys Val Lys Lys Ala Ser Gly Glu Ile Val Ser Ile Asn 50 55 60

Gln Ile Asn Glu Ala His Pro Thr Lys Val Lys Asn Phe Gly Val Trp
65 70 75 80

Val Arg Tyr Asp Ser Arg Ser Gly Thr His Asn Met Tyr Lys Glu Ile 85 90 95

Arg Asp Val Ser Arg Val Ala Ala Val Glu Thr Leu Tyr Gln Asp Met
100 105 110

Ala Ala Arg His Arg Ala Arg Phe Arg Ser Ile His Ile Leu Lys Val 115 120 125



Ala Glu Ile Glu Lys Thr Ala Asp Val Lys Arg Gln Tyr Val Lys Gln 130 135 140

Phe Leu Thr Lys Asp Leu Lys Phe Pro Leu Pro His Arg Val Gln Lys 145 150 155 160

Ser Thr Lys Thr Phe Ser Tyr Lys Arg Pro Ser Thr Phe Tyr 165 170

<210> 267

<211> 932

<212> DNA

<213> Saccharomyces cerevisiae

<400> 267

cttcttacaa gttaatgaat agtatatata tgtaaaaaaa aattctgctt catgtatata 60 cgcacctcgt tattgaggta ttgcgaatgt agcagttgaa taagcaaaga cgaggtttgt 120 acatcttggc atattaaatg gttatttcgg ggtttgtttc ggctcaacgg tgatataaaa 180 agaaatccag tactgtttac cattgtgcat ccgtacattt gatttgttta caacatcttc 240 acatttccat tgtggaatca ttttattttt ttcagaatac cttacccgca agcaaacttt 300 caagcaaact tacaatttgc aaaatttcat cgaattctcc gcaggacata ttataaaagt 360 tatgtcatct cttatcacaa caggtctata agtgttcctt gttcaattgt ttttcccaat 420 tcaaaactct tcacggagca gtttaattat cttactgtcg aagaagtcaa aaactagact 480 atatattatt gagaagaaaa atgtctgacg ttgaagaagt cgttgaagtt caagaagaaa 540 ctgttgttga acaaactgcc gaagttacta tcgaagatgc tttgaaggtt gttttgagaa 600 ccgctttggt tcacgatggt ctagctagag gtttgagaga atctaccaag gctttaacca 660 gaggtgaagc tttattggtt gttttggtca gctctgttac tgaagctaac attatcaagt 720 tggttgaagg tttggctaac gacccagaaa acaaggttcc attgatcaag gttgctgatg 780 ctaagcaatt aggtgaatgg gctggtttgg gtaagatcga ccgtgaaggt aacgccagaa 840 aggttgtcgg tgcctccgtt gttgttgtca agaactgggg tgctgaaact gatgaattgt 900 932 ccatgatcat ggaacacttc tcccaacaat aa

<210> 268

<211> 143

<212> PRT

<213> Saccharomyces cerevisiae

<400> 268

Met Ser Asp Val Glu Glu Val Val Glu Val Glu Glu Glu Thr Val Val

1 5 10 15

Glu Gln Thr Ala Glu Val Thr Ile Glu Asp Ala Leu Lys Val Val Leu
20 25 30

Arg Thr Ala Leu Val His Asp Gly Leu Ala Arg Gly Leu Arg Glu Ser

40

15

Thr Lys Ala Leu Thr Arg Gly Glu Ala Leu Leu Val Val Leu Val Ser 50 55 60 .

Ser Val Thr Glu Ala Asn Ile Ile Lys Leu Val Glu Gly Leu Ala Asn 65 70 75 80

Asp Pro Glu Asn Lys Val Pro Leu Ile Lys Val Ala Asp Ala Lys Gln 85 90 95

Leu Gly Glu Trp Ala Gly Leu Gly Lys Ile Asp Arg Glu Gly Asn Ala 100 105 110

Arg Lys Val Val Gly Ala Ser Val Val Val Val Lys Asn Trp Gly Ala 115 120 125

Glu Thr Asp Glu Leu Ser Met Ile Met Glu His Phe Ser Gln Gln 130 135 140

<210> 269

<211> 800

<212> DNA

<213> Saccharomyces cerevisiae

<400> 269

ggttaggggc caagattacg ttccagcttt cgatgtcgca ccagattggg aatcttacga 60 ctatgcaaag ttggatccaa ctaacgacga cgacaaagaa ttcatcaata acatgtgggc 120 ctgggataag ccagtttccg tcaatggcga accaaaggaa attgttgacg gtaaggtctt 180 aaaataatct cttcgtacta tccttcatgt cgcctttat tataaagtat gctaggtagt 240 tttatctata tcttattat gacgcaatat agggtaacag agttttctg ctctgaaact 300 tccgcagaaa aaaaatcaag ttttcctttt cgtatcttgg attattgtta tataatagat 360 gcatgtatta tatgtatagc agtgatttgc ttatttctt gattctgagg aatcgaagaa 420 gtaacttagc gtttctccg ttgagcagag cagtataata atccttcaaa ttctttaggt 480 tacggggttt tcctgttg atgaccgaag aaactattac tatagattct atttcaaatg 540 ggatactgaa taacctgtta accacattga tccaggacat tgtagctcgg gaaaccactc 600 aacaacaatt gctgaagaca agatatccgg atcttcgcag ttattattc gacccgaacg 660 gatctctcga tattaatgga ctacagaagc aacaagagtc ctctcagtat attcactgtg 720 agaattgtgg cagggatgtg tccgcaaaca gactagcagc tcatttacag agatgtttga 780 gtaggggtgc tagacgttga

<210> 270

<211> 99

<212> PRT

<213> Saccharomyces cerevisiae

```
<400> 270
```

Met Thr Glu Glu Thr Ile Thr Ile Asp Ser Ile Ser Asn Gly Ile Leu
1 5 10 15

Asn Asn Leu Leu Thr Thr Leu Ile Gln Asp Ile Val Ala Arg Glu Thr
20 25 30

Thr Gln Gln Leu Leu Lys Thr Arg Tyr Pro Asp Leu Arg Ser Tyr 35 40 45

Tyr Phe Asp Pro Asn Gly Ser Leu Asp Ile Asn Gly Leu Gln Lys Gln 50 55 60

Gln Glu Ser Ser Gln Tyr Ile His Cys Glu Asn Cys Gly Arg Asp Val 65 70 75 80

Ser Ala Asn Arg Leu Ala Ala His Leu Gln Arg Cys Leu Ser Arg Gly 85 90 95

Ala Arg Arg

<210> 271

<211> 1605

<212> DNA

<213> Saccharomyces cerevisiae

## <400> 271

gtaagcagaa gccggaagaa gacaagaggt tcattcagag aaaacatccg tacattcgag 60 ttctcattga acccatacat ttcaactatt tttacatagt tcgtttttca tgtgtaaaat 120 tgtcatcgac gcgccggagc atcggaaagt tggaacgtgc gcgattgcac caatcccact 180 ggggccgtgc attctgtagg caggaagcca ctggacactc tgcccgttcc cacttggaag 240 attggcgtaa ttccacgctc ctctatcgat tctagcggga aagttatctc tcctggtaag 300 cgttggagga atgccgctac ctaggtaagt ctactgggtg ggaattccag taccgacgtc 360 taggaacatc atgatgctgc agtttctttg aaatttcata tacagtgtta ccaaggataa 420 agtggttaat gcttattcgt cattgaattt ttatatttgc ttaagagaag tgacaaaaga 480 gtgaagacag actatacatc atgaaggtat gatttatgat actattattg aggggcaaca 540 gagaacttta tatgtggaaa aatggcatga aagtttgaaa gtgagaaaga actaaacaga 600 accagacgtc gtaaagattt tagaaatttt ttatggaaag aagcaggaag actaccatat 660 actaattgat gcgtttgtgg cgttattttt aaacaaacag agcgggttta catatactgt 720 tactacaaac cgtgaaattg aagcagttca tatacttgga gtatagtcaa taaaagacaa 780 gctttttcat cttccactta ataatttctc tagagtaata aacttcaccc tattcaaaca 840 ttaattttgc aatagtatac taacaaattt tcttaataac ctgttgaaaa tttaaaatag 900 ttgaacattt cttacccagt taacgggtct caaaagacct tcgaaattga tgatgaacac 960 cgtattcgtg ttttcttcga caagagaatc ggtcaagaag tcgatggtga agccgttggt 1020 gacgaattca agggttaest cttcaagatc tctggtggta acgacaaca aggtttccca 1080 atgaagcaag gtgttttgtt gccaactaga atcaagttgt tgttgaccaa gaacgtttct 1140 tgttacagac caagacgtga tggtgaaaga aagagaaagt ccgtcagagg tgccattgtt 1200 ggtccagatt tggctgtctt ggctttggtc attgtcaaga agggtgagca agaattggaa 1260 ggtctaactg acactactgt tccaaagaga ttgggtccaa agaggctaa caacatcaga 1320 aagttcttcg gtttgtccaa ggaagatgac gttcgtgatt tcgtcatcag aagagaagtc 1380 accaagggtg aaaagactta caccaaggct ccaaagatcc aaagattggt tactcctcaa 1440 agattgcaaa gaagagaca ccaaagagct ttgaaggtca gaaacgctca aggctcaaaga 1500 gaagctgctg ccgaatacgc tcaattgttg gctaagagat tgtctgaaag aaaggctgag 1560 aaggctgaaa tcagaaagag aagagcttct tctttgaagg cttaa 1605

<210> 272

<211> 236

<212> PRT

<213> Saccharomyces cerevisiae

<400> 272

Met Lys Leu Asn Ile Ser Tyr Pro Val Asn Gly Ser Gln Lys Thr Phe 1 5 10 15

Glu Ile Asp Asp Glu His Arg Ile Arg Val Phe Phe Asp Lys Arg Ile 20 25 30

Gly Gln Glu Val Asp Gly Glu Ala Val Gly Asp Glu Phe Lys Gly Tyr 35 40 45

Val Phe Lys Ile Ser Gly Gly Asn Asp Lys Gln Gly Phe Pro Met Lys 50 55 60

Gln Gly Val Leu Leu Pro Thr Arg Ile Lys Leu Leu Leu Thr Lys Asn 65 70 75 80

Val Ser Cys Tyr Arg Pro Arg Arg Asp Gly Glu Arg Lys Arg Lys Ser 85 90 95

Val Arg Gly Ala Ile Val Gly Pro Asp Leu Ala Val Leu Ala Leu Val 100 105 110

Ile Val Lys Lys Gly Glu Gln Glu Leu Glu Gly Leu Thr Asp Thr Thr

Val Pro Lys Arg Leu Gly Pro Lys Arg Ala Asn Asn Ile Arg Lys Phe 130 135 140

Phe Gly Leu Ser Lys Glu Asp Asp Val Arg Asp Phe Val Ile Arg Arg 145 150 155 160



Glu Val Thr Lys Gly Glu Lys Thr Tyr Thr Lys Ala Pro Lys Ile Gln 165 170 175

Arg Leu Val Thr Pro Gln Arg Leu Gln Arg Lys Arg His Gln Arg Ala 180 185 190

Leu Lys Val Arg Asn Ala Gln Ala Gln Arg Glu Ala Ala Ala Glu Tyr
195 200 205

Ala Gln Leu Leu Ala Lys Arg Leu Ser Glu Arg Lys Ala Glu Lys Ala 210 215 220

Glu Ile Arg Lys Arg Arg Ala Ser Ser Leu Lys Ala 225 230 235

<210> 273

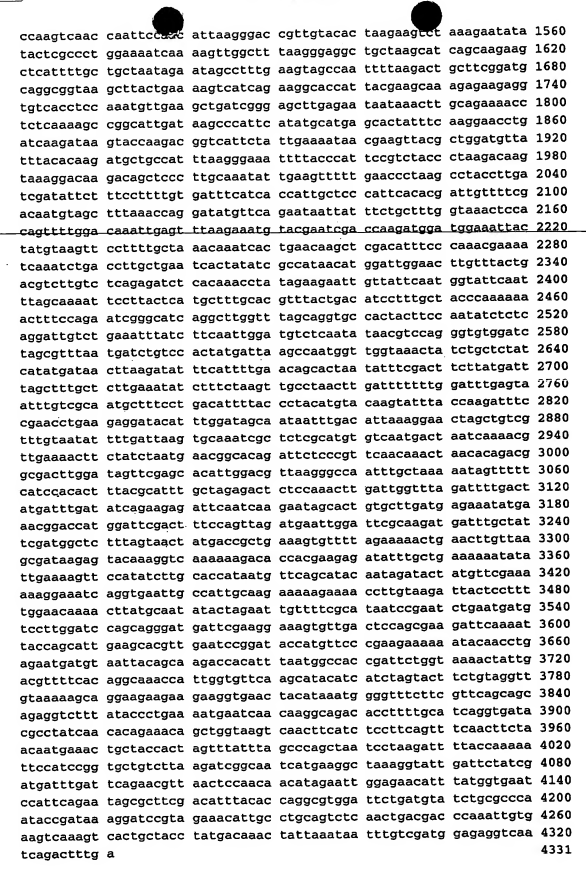
<211> 4331

<212> DNA

<213> Saccharomyces cerevisiae

### <400> 273

gtctcagtga tgcaatcatc atggcacggg cattaattaa ggttaaggaa ccacttctct 60 acgccatact ccaaacaata acaatattag acctagccaa ttgataccca tgagcgtgca 120 gtcattgcca tggattattc atctcttttc gctgttctct ttcttttcaa gtttgtcatc 180 atcatgcctt cacttttgcc tttccatctt tctttttgct gcaaaataaa gggaagaggg 240 gtaaaaacgc aaggaagaac aagaagaaga gggtagtgca agaaaaagaa aagaagaaaa 300 aaaaaaaaaa gtaatCttga taccgtgagc aaataagcta acggaaagcg taagaaagaa 360 gagcgtgttt tgggaaataa caccacagca taaagctaaa attcagttta tataatctat 420 agtagtecta tagaaattge gaataacgga aacaatagte caccaaagca agcataggga 480 gtggagatag catctaggtt atgataacta acaccgagtt cgatgtgccc gtagattggt 540 tgtataaagg taagagtaga cggaagacaa atacgaagcc atcgaggcct tctacgtcgc 600 cagceteete tteatetaeg teetettega aaaaeggaga caatageaeg agtggtaaca 660 ggtctagcaa tgataagcct cgtgcgaggt cgtcatctgt atccaatgca gcactttgta 720 atactgagaa accagatttg aagaggaacg atggtaatac ctctgcatca gacactgata 780 acataceget acttactect attaatageg gtaaceggag egatteeget gacattgata 840 atcoggctac cgtagacgcc atagacctta tagataatga tgacaacggc tcaagtactc 900 aatttgtaag gaaaaaacgt tccacttcta tatccaatgc tgtggtctcg tccaaaccaa 960 gactggccag ttctgccata aacgccaccg catcttcttc tgttggcaag ggaaagcacc 1020 cgccaatttc ttcaccatct aatgctactc tcaagagaag caactcgact agtggagaaa 1080 agactaaaag gtcgattttt ggatctttgt ttagcaaacg gtctacctct tcttcagctt 1140 ccactgccaa gaaaccatta cctgttgtta atactagcac gactgagaat gagtctggtg 1200 gcattaaagc agttgcgact cctgatccac gggtgaagga gatatcatca ccaatgagag 1260 taaaagacct atctacagtt tcattgaaaa gggtctcgtt tgccgtcgat aaattcgaat 1380 ccgatcctcc ccaacaactt ccttcaagaa ccccaaaaaa gggaaacatc cttatccctg 1440 atgacatgat aagcgaggtt ccttctattt ccgtgggtat ttccagtagc aaccaatcgg 1500



<210> 274

<211> 1276

<212> PRT

<213> Saccharomyces cerevisiae

<400> 274

Met Ile Thr Asn Thr Glu Phe Asp Val Pro Val Asp Trp Leu Tyr Lys

1 5 10 15

Gly Lys Ser Arg Arg Lys Thr Asn Thr Lys Pro Ser Arg Pro Ser Thr
20 25 30

Ser Pro Ala Ser Ser Ser Ser Thr Ser Ser Ser Lys Asn Gly Asp Asn 35 40 45

Ser Thr Ser Gly Asn Arg Ser Ser Asn Asp Lys Pro Arg Ala Arg Ser 50 55 60

Ser Ser Val Ser Asn Ala Ala Leu Cys Asn Thr Glu Lys Pro Asp Leu 65 70 75 80

Lys Arg Asn Asp Gly Asn Thr Ser Ala Ser Asp Thr Asp Asn Ile Pro 85 90 95

Leu Leu Thr Pro Ile Asn Ser Gly Asn Arg Ser Asp Ser Ala Asp Ile 100 105 110

Asp Asn Pro Ala Thr Val Asp Ala Ile Asp Leu Ile Asp Asn Asp Asp 115 120 125

Asn Gly Ser Ser Thr Gln Phe Val Arg Lys Lys Arg Ser Thr Ser Ile 130 135 140

Asn Ala Thr Ala Ser Ser Ser Val Gly Lys Gly Lys His Pro Pro Ile 165 170 175

Ser Ser Pro Ser Asn Ala Thr Leu Lys Arg Ser Asn Ser Thr Ser Gly
180 185 190

Glu Lys Thr Lys Arg Ser Ile Phe Gly Ser Leu Phe Ser Lys Arg Ser 195 200 205

Thr Ser Ser Ser Ala Ser Thr Ala Lys Lys Pro Leu Pro Val Val Asn

215 220 210 Thr Ser Thr Thr Glu Asn Glu Ser Gly Gly Ile Lys Ala Val Ala Thr 235 230 225 Pro Asp Pro Arg Val Lys Glu Ile Ser Ser Pro Met Arg Gly Val Ala 250 Pro Thr Ala Ser Lys Pro Gln Thr Pro Ile Leu Pro Ser Pro Ala Leu 265 Ala Val Lys Asp Leu Ser Thr Val Ser Leu Lys Arg Val Ser Phe Ala 285 280 Val Asp Lys Phe Glu Ser Asp Pro Pro Gln Gln Leu Pro Ser Arg Thr 300 295 290 Pro Lys Lys Gly Asn Ile Leu Ile Pro Asp Asp Met Ile Ser Glu Val 315 310 305 Pro Ser Ile Ser Val Gly Ile Ser Ser Ser Asn Gln Ser Ala Lys Ser 330 325 Thr Asn Ser Asn Ile Lys Gly Pro Leu Tyr Thr Lys Lys Ser Lys Glu 345 340 Tyr Ile Leu Ala Leu Glu Asn Gln Lys Leu Ala Leu Arg Glu Ala Ala 360 Lys His Gln Gln Glu Ala His Phe Ala Ala Asn Arg Ile Ala Phe Glu 380 375 Val Ala Asn Phe Lys Thr Ala Ser Asp Ala Gly Gly Lys Leu Thr Glu 400 395 390 385 Lys Ser Ser Glu Gly Thr Ile Thr Lys Gln Arg Glu Glu Val Ser Pro 410 405 Pro Asn Val Glu Ala Asp Arg Glu Leu Glu Asn Asn Lys Leu Ala Glu

Asn Leu Ser Lys Ala Gly Ile Asp Lys Pro Ile His Met His Glu His 435 440 445

420

425

Tyr Phe Lys Glu Pro Asp Gln Asp Lys Tyr Gln Asp Gly His Ser Ile 450 455 460

Glu Asn Asn Glu Val Thr Leu Asp Val Ile Tyr Thr Arg Cys Cys His

Leu Arg Glu Ile Leu Pro Ile Pro Ser Thr Leu Arg Gln Val Lys Asp Lys Thr Ala Pro Leu Gln Ile Leu Lys Phe Leu Asn Pro Lys Pro Thr Leu Ile Asp Ile Leu Ser Phe Cys Asp Phe Ile Thr Ile Ala Pro Ile His Thr Ile Val Phe Asp Asn Val Ala Leu Asn Gln Asp Met Phe Arg Ile Ile Ile Ser Ala Leu Val Asn Ser Thr Val Leu Asp Lys Leu Ser Leu Arg Asn Val Arg Ile Asp Gln Asp Gly Trp Lys Leu Leu Cys Lys Phe Leu Leu Leu Asn Lys Ser Leu Asn Lys Leu Asp Ile Ser Gln Thr Lys Ile Lys Ser Asp Leu Ala Glu Ser Leu Tyr Arg His Asn Met Asp Trp Asn Leu Phe Thr Asp Val Leu Ser Gln Arg Ser His Lys Pro Ile Glu Glu Leu Leu Phe Asn Gly Ile Gln Phe Ser Lys Ile Pro Tyr Ser Cys Phe Ala Arg Leu Leu Thr Ser Phe Ala Thr Gln Lys Asn Phe Pro Glu Ser Gly Ile Arg Leu Gly Leu Ala Gly Ala Thr Thr Ser Asn Ile Ser Gln Asp Cys Leu Lys Phe Ile Phe Asn Trp Met Ser Gln Tyr Asn Val Gln Gly Val Asp Leu Ala Phe Asn Asp Leu Ser Thr Met Ile Lys 

Phe Ile Leu Asn Ser Thr Asn Ile Ser Thr Ser Tyr Asp Leu Ala Leu

Pro Met Val Gly Lys Leu Ser Ala Leu Ser Tyr Asp Asn Leu Arg Tyr

730

735

Leu Leu Lys Tyr Leu Ser Lys Leu Pro Asn Leu Ile Phe Leu Asp Leu 740 745 750

Ser Asn Leu Ser Gln Cys Phe Pro Asp Ile Leu Pro Tyr Met Tyr Lys
755 760 765

Tyr Leu Pro Arg Phe Pro Asn Leu Lys Arg Ile His Leu Asp Ser Asn 770 775 780

Ash Leu Thr Leu Lys Glu Leu Ala Val Val Cys Ash Ile Leu Ile Lys
785 790 795 800

Cys Lys Ser Leu Ser His Val Ser Met Thr Asn Gln Asn Val Glu Asn 805 810 815

Phe Tyr Leu Met Asn Gly Thr Asp Ser Pro Val Gln Gln Thr Asn Thr 820 825 830

Asp Gly Asp Leu Asp Ser Ser Ser Thr Leu Asp Val Lys Gly Gln Phe 835 840 845

Ala Lys Asn Ser Phe Ser Ser Thr Leu Tyr Ala Phe Ala Arg Asp Ser .850 855 860

Pro Asn Leu Ile Gly Leu Asp Phe Asp Tyr Asp Leu Ile Ser Glu Glu 865 870 875 880

Ile Gln Ser Arg Ile Ala Leu Cys Leu Met Arg Asn Met Lys Arg Thr 885 890 895

Met Asp Ser Thr Phe Gln Leu Asp Glu Leu Asp Ser Gln Asp Asp Leu 900 905 910

Leu Phe Asp Gly Ser Leu Val Thr Met Thr Ala Glu Ser Val Leu Glu 915 920 925

Lys Leu Asn Leu Leu Ser Asp Lys Ser Thr Lys Val Lys Lys Asp Thr 930 935 940

Thr Lys Arg Tyr Leu Leu Lys Lys Tyr Ile Glu Lys Phe His Ile Leu 945 950 955 960

His His Asn Val Gln His Thr Ile Asp Thr Met Phe Glu Lys Arg Lys 965 970 975

Ser Gly Glu Leu Pro Leu Gln Glu Lys Glu Asn Leu Val Arg Leu Leu

985

990

Leu Leu Glu Gln Asn Leu Cys Asn Ile Leu Glu Leu Phe Ser His Asn 995 1000 1005

Pro Asn Leu Asn Asp Val Leu Gly Ser Ser Arg Asp Asp Ser Lys Glu 1010 1015 1020

Ser Val Asp Ser Ser Glu Asp Ser Lys Leu Pro Ala Leu Lys His Val 1025 1030 1035 1040

Glu Ser Gly Tyr His Val Pro Glu Glu Lys Ile Gln Pro Glu Asn Asp 1045 1050 1055

Val Ile Thr Ala Arg Pro His Leu Met Ala Thr Asp Ser Gly Lys Thr 1060 1065 1070

Ile Asp Val Phe Thr Gly Lys Pro Leu Val Phe Lys His Thr Ser Ser 1075 1080 1085

Ser Thr Ser Val Gly Cys Lys Lys Gln Glu Glu Glu Glu Glu Glu Leu 1090 1095 1100

His Lys Trp Gly Phe Phe Val Gln Gln Gln Arg Ser Leu Tyr Pro Glu 1105 1110 1115 1120

Asn Glu Ser Thr Arg Gln Thr Pro Phe Ala Ser Gly Asp Thr Pro Ile 1125 1130 1135

Asn Thr Glu Thr Ala Gly Lys Ser Thr Ser Ser Pro Ser Val Ser Thr 1140 1145 1150

Ser Asn Asn Glu Thr Ala Thr Thr Ser Leu Phe Ser Pro Ala Asn Pro 1155 1160 1165

Lys Ile Leu Pro Lys Ile Pro Ser Gly Ala Val Leu Arg Ser Ala Ile 1170 1175 1180

Met Lys Ala Lys Gly Ile Asp Ser Ile Asp Asp Leu Ile Gln Asn Val 1185 1190 1195 1200

Asn Ser Asn Asn Ile Glu Leu Glu Asn Ile Tyr Gly Glu Ser Ile Gln
1205 1210 1215

Asn Ser Ala Ser Thr Phe Thr Pro Gly Val Asp Ser Asp Val Ser Ala 1220 1225 1230

Pro Asn Thr Asp Lys Gly Ser Val Glu Thr Leu Pro Ala Val Ser Thr

1240



Asp Asp Pro Asn Cys Glu Val Lys Val Thr Ala Thr Tyr Asp Lys Leu 1250 1255 1260

Leu Asn Asn Leu Ser Met Glu Arg Ser Il Arg Leu 1265 1270 1275

<210> 275

<211> 1262

<212> DNA

<213> Saccharomyces cerevisiae

<400> 275

aaacaaagcg atttgagtat tacgttaaga tctccatttg ctatgctaaa tcctgcctac 60 tcaaacagca ttatatcgct accaacaggt gtggtcaaga ggtcattgac tcttcctgtc 120 ggtatgaaaa tttaaaaata tataataaca taagtttgca ttcattaata ttaatataaa 180 traacqtatc aaccataaga accataatca taataattgt ttaccttaat ggcgatatcg 240 ccttttatca ccatcgacga taacaaacgc ctattctgtc taccgtttct ttttttcgct 300 cgcaatatca gaatcgttta tcccaaaggg tagttatagc aactatttaa ataacttaga 360 tgtttcttta acttattccg tcatacgtac acaaacaccc gctagatata gcactccttg 420 ctgtcgaaca ttataaaggt gcttttaaaa ctactaatcg tatattcagc aggtcagaac 480 gcaaagtcgg acgataaact atgttgaagc tagctcgtcc atttattccg cctttatcaa 540 ggaacaatgc catttcttca ggaatagttc tcacttctag aagatttcag tcttccttta 600 cgttcttaag taaccagtct ttactatcta aaaatcaaat gaaatccaaa agaaaaaagg 660 gcagtaaaaa ggcagcgtac catcgtcaac ccccggaaca cgaacatact gcaccactta 720 taaagcaaaa caagacaatc acaaagaaag aacatagcga tgtgagaggt tctcatttaa 780 aaaagaaaag aagcgatttt tcgtggctac caagagtacc atccacctca catttaaagc 840 agagtgacat gaccacaaat gtactctatt ctggatatag acccttattc atcaatccca 900 atgacccgaa gctaaaggaa gacaccggaa gtacgttata cgaatttgcg atgaagcttg 960 aagatttaaa tgaacctcta tcaccatgga tttcctctgc cactggactt gaattctttt 1020 cagagtggga gaatatacct agtgaactac tgaaaaattt gaagcctttt catccaccta 1080 aggaaaaatc aatgaacaca aatgagctta ttcatgtaag tgctaagaga aatacattag 1140 gaaaggggag aaaaaagtca gtagtgacgt tgctccaaat gaagaaaaaa cttgagggat 1260 1262 ga

<210> 276

<211> 253

<212> PRT

<213> Saccharomyces cerevisiae

<400> 276

Met Leu Lys Leu Ala Arg Pro Phe Ile Pro Pro Leu Ser Arg Asn Asn 1 5 10 15

	Ala	IIe	ser	20	GIĀ	11e	Val	Leu	25	ser	Arg	Arg	Pne	30	ser	ser
	Phe	Thr	Phe 35	Leu	Ser	Asn	Gln	Ser 40	Leu	Leu	Ser	Lys	Asn 45	Gln	Met	Lys
	Ser	Lys 50	Arg	Lys	Lys	Gly	Ser 55	Lys	Lys	Ala	Ala	Tyr 60	His	Arg	Gln	Pro
	Pro	Glu	His	Glu	His	Thr	Ala	Pro	Leu	Ile	Lys	Gln	Asn	Lys	Thr	Ile
_	65					70					75					80
	Thr	Lys	Lys	Glu	His 85	Ser	Asp	Val	Arg	Gly 90	Ser	His	Leu	Lys	Lys 95	Lys
	Arg	Ser	Asp	Phe 100	Ser	Trp	Leu	Pro	Arg 105	Val	Pro	Ser	Thr	Ser 110	His	Leu
	Lys	Gln	Ser 115	Asp	Met	Thr	Thr	Asn 120	Val	Leu	Tyr	Ser	Gly 125	Tyr	Arg	Pro
	Leu	Phe 130	Ile	Asn	Pro	Asn	Asp 135	Pro	Lys	Leu	Lys	Glu 140	Asp	Thr	Gly	Ser
	Thr 145	Leu	Tyr	Glu	Phe	Ala 150	Met	Lys	Leu	Glu	Asp 155	Leu	Asn	Glu	Pro	Leu 160
	Ser	Pro	Trp	Ile	Ser 165	Ser	Ala	Thr	Gly	Leu 170	Glu	Phe	Phe	Ser	Glu 175	Trp
	Glu	Asn	Ile	Pro 180	Ser	Glu	Leu	Leu	Lys 185	Asn	Leu	Lys	Pro	Phe 190	His	Pro
	Pro	Lys	Glu 195	Lys	Ser	Met	Asn	Thr 200	Asn	Glu	Leu	Ile	His 205	Val	Ser	Ala
	Lys	Arg 210	Asn	Thr	Leu	Val	Asp 215	Asn	Lys	Thr	Ser	Glu 220	Thr	Leu	Gln	Arg
	Lys 225	Met	Asp	Glu	Phe	Ser 230	Lys	Arg	Arg	Gly	Lys 235	Gly	Arg	Lys	Lys	Ser 240
	Val	Val	Thr	Leu	Leu 245	Gln	Met	Lys	Lys	Lys 250	Leu	Glu	Gly			

<210> 277
<211> 1940
<212> DNA

<213> Saccharomyces cerevisiae

```
<400> 277
acgttatttt caaaacacat acgaaatcgc tggactttct cactaaaggc ttccggagcc 60
tgcggttctt ccatccccat ctcttccaca tataccgggg gttctgatgg cattcctata 120
gcttgctcaa agtcctgcga gtttagctca tcttgtgagt tatttgctgg ttcttcaaca 180
ttcggaagag tagtgtagcc cctagtattc accatatcct ttgtatactc atatgcgaac 240
ttgtctaata caatgtatac caaaaataac agcttgaaaa aaatctcagg gtcttttcat 300
tgaataatet cattetgeta ttttaagttt ttegtttgea accegagact gtcgagetag 360
aaaatttcat tgatacgatt taaaaaataat cgatgccagt aatagcctca aaaaacttat 420
gcaaaaatac gagatactat aaactacagc ttagctaact ctaacattat tatataaaac 480
aatgggcttc aatatagcgt atgtctagct cacagcatgt gttccaaata cattaaagaa 540
gatctctttt gttgttgata ctaaccagta aagttgagag ttataacaat gaaaatagga 600
tgctgtgcga ctttttttat ccacagttag gtggagtcga attccatata tatcatttat 660
cgcagaaact aatcgatttg ggccattctg tcgtcattat aactcacgct tacaaagatc 720
gagtcggcgt acgacatett accaacggte taaaggteta teacgtacea ttttttgtga 780
ttttcagaga aaccactttc cccactgttt tttcaacatt tccaataata aggaatattc 840
ttctcagaga gcagatccaa attgttcatt ctcatggtag cgcttccacg ttcgctcacg 900
agggaattet teatgetaat actatgggat tgagaactgt gtteaeggae catteaetet 960
acggttttaa taacttaacg tcgatttggg tgaataagtt gctaacattt accttgacaa 1020
acatagatcg ggttatatgt gtttctaata catgcaaaga aaatatgatt gttagaacag 1080
aattaagtcc tgatataatc tcagtaattc ccaacgcagt ggtgagcgaa gatttcaaac 1140
caagggatcc tactggtggc accaagagaa aacaaagtag ggataagata gtgatcgtgg 1200
tcatcggaag gctctttcca aacaaagggt ccgatttact tactcgcata attccgaaag 1260
tttgttcctc acatgaagat gtcgaattta tagtagcggg cgatggtcca aagttcatag 1320
attttcaaca aatgattgaa agtcatagac tacaaaaacg tgtgcaactc ttaggctctg 1380
ttccacatga gaaagtcagg gatgtattat gtcaaggtga catatattta cacgctagtt 1440
taacagaagc atttggtaca attctagttg aggccgcatc ttgtaatttg ctaattgtaa 1500
cgacacaagt cggaggaatt cccgaagtgt taccaaatga gatgactgtt tatgcagaac 1560
agacatccgt ttctgacctt gttcaagcaa caaataaagc tatcaatatc ataagaagta 1620
aagetttgga cactteetet ttteatgata gegtgtetaa aatgtaegae tggatggaeg 1680
tagecaaaag gacagtagag atatatacta atatatette taettettee getgatgata 1740
aagattggat gaaaatggta gcaaatcttt acaaaagaga tggaatctgg gctaaacatc 1800
tttatctgtt atgtggaatt gtagagtaca tgcttttttt cctcttagag tggctatacc 1860
ccagggatga aatcgatcta gctccaaaat ggcccaagaa aacagtgtct aacgagacga 1920
                                                                   1940
aggaagcaag agaaacttaa
```

```
<210> 278
```

# <400> 278

Met Ser Ser Ser His Lys Val Glu Ser Tyr Asn Asn Glu Asn Arg Met

<sup>&</sup>lt;211> 461

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Saccharomyces cerevisiae

1	5	10

Leu Cys Asp Phe Phe Tyr Pro Gln Leu Gly Gly Val Glu Phe His Ile 20 25 30

Tyr His Leu Ser Gln Lys Leu Ile Asp Leu Gly His Ser Val Val Ile
35 40 45

Ile Thr His Ala Tyr Lys Asp Arg Val Gly Val Arg His Leu Thr Asn 50 55 60

Gly Leu Lys Val Tyr His Val Pro Phe Phe Val Ile Phe Arg Glu Thr
65 70 75 80

Thr Phe Pro Thr Val Phe Ser Thr Phe Pro Ile Ile Arg Asn Ile Leu 85 90 95

Leu Arg Glu Gln Ile Gln Ile Val His Ser His Gly Ser Ala Ser Thr
100 105 110

Phe Ala His Glu Gly Ile Leu His Ala Asn Thr Met Gly Leu Arg Thr 115 120 125

Val Phe Thr Asp His Ser Leu Tyr Gly Phe Asn Asn Leu Thr Ser Ile 130 135 140

Trp Val Asn Lys Leu Leu Thr Phe Thr Leu Thr Asn Ile Asp Arg Val 145 150 155 160

Ile Cys Val Ser Asn Thr Cys Lys Glu Asn Met Ile Val Arg Thr Glu
165 170 175

Leu Ser Pro Asp Ile Ile Ser Val Ile Pro Asn Ala Val Val Ser Glu 180 185 190

Asp Phe Lys Pro Arg Asp Pro Thr Gly Gly Thr Lys Arg Lys Gln Ser 195 200 205

Arg Asp Lys Ile Val Ile Val Val Ile Gly Arg Leu Phe Pro Asn Lys 210 225 220

Gly Ser Asp Leu Leu Thr Arg Ile Ile Pro Lys Val Cys Ser Ser His 225 230 235 240

Glu Asp Val Glu Phe Ile Val Ala Gly Asp Gly Pro Lys Phe Ile Asp 245 250 255

Phe Gln Gln Met Ile Glu Ser His Arg Leu Gln Lys Arg Val Gln Leu

260

Leu Gly Ser Val Pro His Glu Lys Val Arg Asp Val Leu Cys Gln Gly 280

Asp Ile Tyr Leu His Ala Ser Leu Thr Glu Ala Phe Gly Thr Ile Leu 295

Val Glu Ala Ala Ser Cys Asn Leu Leu Ile Val Thr Thr Gln Val Gly 315 310

Gly Ile Pro Glu Val Leu Pro Asn Glu Met Thr Val Tyr Ala Glu Gln 335

330 325

Thr Ser Val Ser Asp Leu Val Gln Ala Thr Asn Lys Ala Ile Asn Ile 345 340

Ile Arg Ser Lys Ala Leu Asp Thr Ser Ser Phe His Asp Ser Val Ser 360 355

Lys Met Tyr Asp Trp Met Asp Val Ala Lys Arg Thr Val Glu Ile Tyr 380 375 370

Thr Asn Ile Ser Ser Thr Ser Ser Ala Asp Asp Lys Asp Trp Met Lys 395 390 385

Met Val Ala Asn Leu Tyr Lys Arg Asp Gly Ile Trp Ala Lys His Leu 410 405

Tyr Leu Leu Cys Gly Ile Val Glu Tyr Met Leu Phe Phe Leu Leu Glu 425 420

Trp Leu Tyr Pro Arg Asp Glu Ile Asp Leu Ala Pro Lys Trp Pro Lys 445 435 440

Lys Thr Val Ser Asn Glu Thr Lys Glu Ala Arg Glu Thr 460 455 450

<210> 279

<211> 2900

<212> DNA

<213> Saccharomyces cerevisiae

<400> 279

ctcaatggga attgagtgaa gatcagtctg atcgagagga atttgtacga ttcgtggaaa 60 accagcattt cgtagaaaaa gttgatacga tttacaacgg ttataatgaa agtctatcaa 120

```
tgatggacga cctgactaga gagttactac tatgggagaa aaaatattca aataacacta 180
atgccattca ataaacgcaa aacactgcaa tattattctc aaccaaagta taactgtaat 240
gaggcgaaca aacacatcta tacatatata tacatctata tggatataaa aacgactaat 300
tcaacgttgt ttttatcaac cgagcttact cttgtacggg taaccgcaag gatagctagt 360
tgcggatggt atagcgattt ggctggcacg atgattaagg aatccaaaca tctaatggac 420
tagcacattc tatcgattta cgggtcaggt aaacatagat attgggatat atcatatatc 480
cttactgagt aactataatt atggttcatc gaggaaggac tttgaagtca gacactgatg 540
taacatetet taatgegtea acagtateae accagteaaa gecatttaga cagttttega 600
ctaggtcgag agcaaagagt aacgcaagct tcaaaggttt gcgtagagtt ttaacacatg 660
atggcaccct ggataatgat tattttaata agcacaacgt ttctcagaaa tgcaagagtt 720
ctgatgcact tttcagaaag cgaacgatta gtgggttgaa tatgacagct ttaacaagag 780
taaagtecaa teaaggaaaa agateageat eetileatag teeggtgeat aataegetge 840
tcagtccaaa gaacagcagt cattctaata ctggaactgc tggtttcggc ctgaaaccac 900
gaagaagtaa aagtacccaa tetgttetga gtettegaga tgegcaagaa tetaaaaaga 960
gtgaatctac tactgacgag gaggtggaat gtttttcgga agacaacatt gaagatggaa 1020
aggtgaataa tgataaagta atagccgagc atgttatgcc tgaagaaaaa aagaatgtgc 1080
agcaattaaa tcagaatgaa ttacaatccc cggattcaat agatgaacaa gaagaagata 1140
aatcaggtac tgatggaaag gaaaatcata gagctgtatc cttaccatta cctcatttat 1200
cttccaataa ctatttcgga gaatcaagcc attctataga acatcagaaa gatggagaaa 1260
catctccaag ctcaattgaa acaaaactga atgcaacaag tgtaatcaat gaagaggggc 1320
aatcaaaggt gacgaaggaa gctgatattg atgacttgtc cagccattct caaaatttga 1380
gggcctcatt ggttaaagcg ggcgataata tatcagaagc accatatgat aaagaaaaaa 1440
aaattettga tgttggtaat acettagetg cacataaaag taatcaaaaa ccaagteatt 1500
cagatgaaca gtttgatcag gaagatcaca ttgatgcccc taggagtaat tcatcaagaa 1560
aaagcgactc gagctttatg tctcttagga gacaaagttc taaacaacac aaattattaa 1620
acgaagaaga agatctaatc aagcctgatg atatttcttc cgctggtacc aaggatattg 1680
aagggcatag cttactggaa aattatgcgc ctaatatgat tctctcccag tcgactggag 1740
ttgaacgtag atttgaaaat tcatcatcca tccaaaattc gcttgggaat gaaattcatg 1800
actcgggtga gcatatggct tcaggtgata cttttaatga actggatgat ggcaaattgc 1860
gcaagagcaa gaaaaatggt ggaagatctc aacttggcca aaatataccg aactctcagt 1920
ctactttccc caccattgct aacatcggta gtaaagataa taatgtacca cagcacaact 1980
tttcgacctc catatcgagt ttaaccaata atttgaggag agctgctcct gaaagcttcc 2040
atggttcaag aatgaataat atttttcaca agaaaggtaa tcagaatcta cttctgagat 2100
ccaacgatct caacaaaaat tctgcagccc cggcctctcc attgtccaac gaacatatta 2160
catctagtac gaactccggt agcgatgcaa acagacaatc caactcaggt gccaaattta 2220
atagettege ceagtteett aaateagatg ggattgatge agaateaaga acacaaagaa 2280
aattatggtt gcagagggag aattctatta tggacttaag ttcacaaaat gacggtagtg 2340
actctatctt tatggcagga aacattgatg cgaaaaggga gtttgagaga atatcccatg 2400
aatactctaa tgtaaaaaga ttttacaacc cattagatga agcattgttg agagtacaac 2460
ctataataac gggaaatgca aataatatca ggaaaaaaag ccataacgat gctcagtcaa 2520
tcgcacattc tagcagtgat acagatcata aggatgagga cgatttgctc tttactaact 2580
atgacaaaaa atttgatgat ctttatccac atcttgcaag tgcaaagatt caggcagtgt 2640
tgtccggtat atggaaaagc gaaagttact tatttaacaa ggatgttaat ccaatcaaca 2700
agaataggac aacgagtaca aaccacagcg ttggccacac tgcttcacag aatgcacgta 2760
acttgctgag gggcccgatg ggttccagca cgactttgca ccaccaacgc gtcattaact 2820
ctctgcagcc gactacgagg gcagtgaatc gcaggatgga aaatgtgggc tacatgcata 2880
                                                                  2900
cacagccaca acaaaggtga
```

<210> 280 <211> 799

<212> PRT

<213> Saccharomyces cerevisiae

<400> 280

Met Val His Arg Gly Arg Thr Leu Lys Ser Asp Thr Asp Val Thr Ser 1 5 10 15

Leu Asn Ala Ser Thr Val Ser His Gln Ser Lys Pro Phe Arg Gln Phe

20 25 3

Ser Thr Arg Ser Arg Ala Lys Ser Asn Ala Ser Phe Lys Gly Leu Arg 35 40 45

Arg Val Leu Thr His Asp Gly Thr Leu Asp Asn Asp Tyr Phe Asn Lys 50 55 60

His Asn Val Ser Gln Lys Cys Lys Ser Ser Asp Ala Leu Phe Arg Lys
65 70 75 80

Arg Thr Ile Ser Gly Leu Asn Met Thr Ala Leu Thr Arg Val Lys Ser 85 90 95

Asn Gln Gly Lys Arg Ser Ala Ser Phe His Ser Pro Val His Asn Thr 100 105 110

Leu Leu Ser Pro Lys Asn Ser Ser His Ser Asn Thr Gly Thr Ala Gly
115 120 125

Phe Gly Leu Lys Pro Arg Arg Ser Lys Ser Thr Gln Ser Val Leu Ser 130 135 140

Leu Arg Asp Ala Gln Glu Ser Lys Lys Ser Glu Ser Thr Thr Asp Glu 145 150 155 160

Glu Val Glu Cys Phe Ser Glu Asp Asn Ile Glu Asp Gly Lys Val Asn 165 170 175

Asn Asp Lys Val Ile Ala Glu His Val Met Pro Glu Glu Lys Lys Asn 180 185 190

Val Gln Gln Leu Asn Gln Asn Glu Leu Gln Ser Pro Asp Ser Ile Asp 195 200 205

Glu Gln Glu Glu Asp Lys Ser Gly Thr Asp Gly Lys Glu Asn His Arg 210 215 220

Ala 225	Val	Ser	Leu	Pro	Leu 230	Pro	His	Leu	Ser	Ser 235	Asn	Asn	Tyr	Phe	Gly 240
Glu	Ser	Ser	His	Ser 245	Ile	Glu	His	Gln	Lys 250	Asp	Gly	Glu	Thr	Ser 255	Pro
Ser	Ser	Ile	Glu 260	Thr	Lys	Leu	Asn	Ala 265	Thr	Ser	Val	Ile	Asn 270	Glu	Glu
Gly	Gln	Ser 275	_			-			_		Asp	_		Ser	Ser
		213					200					<del>-203</del> -			
His	Ser 290	Gln	Asn	Leu	Arg	Ala 295	Ser	Leu	Val	Lys	Ala 300	Gly	Asp	Asn	Ile
Ser	Glu	Ala	Pro	Tyr	Asp	Lys	Glu	Lys	Lys	Ile	Leu	Asp	Val	Gly	Asn
305					310					315					320
Thr	Leu	Ala	Ala	His 325	Lys	Ser	Asn	Gln	Lys 330	Pro	Ser	His	Ser	Asp 335	Glu
Gln	Phe	Asp	Gln 340	Glu	Asp	His	Ile	Asp 345	Ala	Pro	Arg	Ser	Asn 350	Ser	Ser
Arg	Lys	Ser 355	Asp	Ser	Ser	Phe	Met 360	Ser	Leu	Arg	Arg	Gln 365	Ser	Ser	Lys
Gln	His 370	Lys	Leu	Leu	Asn	Glu 375	Glu	Glu	Asp	Leu	Ile 380	Lys	Pro	Asp	Asp
Ile 385	Ser	Ser	Ala	Gly	Thr 390	Lys	Asp	Ile	Glu	Gly 395	His	Ser	Leu	Leu	Glu 400
Asn	Tyr	Ala	Pro	Asn 405	Met	Ile	Leu	Ser	Gln 410	Ser	Thr	Gly	Val	Glu 415	Arg
Arg	Phe	Glu	Asn 420	Ser	Ser	Ser	Ile	Gln 425	Asn	Ser	Leu	Gly	Asn 430	Glu	Ile
His	Asp	Ser 435	Gly	Glu	His	Met	Ala 440	Ser	Gly	Asp	Thr	Phe 445	Asn	Glu	Leu
Asp	Asp 450	Gly	Lys	Leu	Arg	Lys 455	Ser	Lys	Lys	Asn	Gly 460	Gly	Arg	Ser	Gln
Leu 465	Gly	Gln	Asn	Ile	Pro 470	Asn	Ser	Gln	Ser	Thr 475	Phe	Pro	Thr	Ile	Ala 480

Asn	Ile	Gly	Ser	Lys 485	Asp	Asn	Asn	Val	Pro 490	Gln	His	Asn	Phe	Ser 495	Thr
Ser	Ile	Ser	Ser 500	Leu	Thr	Asn	Asn	Leu 505	Arg	Arg	Ala	Ala	Pro 510	Glu	Ser
Phe	His	Gly 515	Ser	Arg	Met	Asn	Asn 520	Ile	Phe	His	Lys	Lys 525	Gly	Asn	Gln
Asn	Leu 530	Leu	Leu	Arg	Ser	Asn 535	Asp	Leu	Asn	Lys	Asn -540-	Ser	Ala	Ala	Pro
	330														
Ala 545	Ser	Pro	Leu	Ser	Asn 550	Glu	His	Ile	Thr	Ser 555	Ser	Thr	Asn	Ser	Gly 560
Ser	Asp	Ala	Asn	Arg 565	Gln	Ser	Asn	Ser	Gly 570	Ala	Lys	Phe	Asn	Ser 575	Phe
Ala	GÌn	Phe	Leu 580	Lys	Ser	Asp	Gly	Ile 585	Ąsp	Ala	Glu	Ser	Arg 590	Thr	Gln
Arg	Lys	Leu 595	Trp	Leu	Gln	Arg	Glu 600	Asn	Ser	Ile	Met	Asp 605	Leu	Ser	Ser
Gln	Asn 610	Asp	Gly	Ser	Asp	Ser 615	Ile	Phe	Met	Ala	Gly 620	Asn	Ile	Asp	Ala
Lys 625	Arg	Glu	Phe	Glu	Arg 630	Ile	Ser	His	Glu	Tyr 635	Ser	Asn	Val	Lys	Arg 640
Phe	Tyr	Asn	Pro	Leu 645	Asp	Glu	Ala	Leu	Leu 650	Arg	Val	Gln	Pro	Ile 655	Ile
Thr	Gly		Ala 660		Asn									Ala	Gln
Ser	Ile	Ala 675	His	Ser	Ser	Ser	Asp 680	Thr	Asp	His	Lys	Asp 685	Glu	Asp	Asp
Leu	Leu 690	Phe	Thr	Asn	Tyr	Asp 695	Lys	Lys	Phe	Asp	Asp 700	Leu	Tyr	Pro	His
Leu 705	Ala	Ser	Ala	Lys	Ile 710	Gln	Ala	Val	Leu	Ser 715	Gly	Ile	Trp	Lys	Ser 720
Glu	Ser	Tyr	Leu	Phe 725	Asn	Lys	Asp	Val	Asn 730	Pro	Ile	Asn	Lys	Asn 735	Arg

```
Thr Thr Ser Thr Asn His Ser Val Gly His Thr Ala Ser Gln Asn Ala
740 745 750
```

Arg Asn Leu Leu Arg Gly Pro Met Gly Ser Ser Thr Thr Leu His His 755 760 765

Gln Arg Val Ile Asn Ser Leu Gln Pro Thr Thr Arg Ala Val Asn Arg 770 775 780

Arg Met Glu Asn Val Gly Tyr Met His Thr Gln Pro Gln Gln Arg
785 790 795

<210> 281 <211> 1212 <212> DNA <213> Saccharomyces cerevisiae

### <400> 281

tgatacaaga caggcattag gaaacacatg agttttgcat atgtattatg caggttcatt 60 tgttccttcc cttatttctt ttcatagtat ttatttttat ttatttattt attttttt 120 ggattttatt ttatttcttt tttaatgcta agaaagtaat tccgcataat taaacgtgtg 180 ctggcctcga taggtaccta tagtatacag aagcttacga aaagctcctg caggatggca 240 cttctaaaat tcgcgctcaa catggccgta ttgtacatta tatcgttcta tcattatatc 300 gtatacgccc gcattacccg acaactccgt ctgcaacgcg ttgaccagaa aactcgaaca 360 agagatcgca taaaaaacca aaaggaaacg aattacttgt caaatagtta ttgtaatgga 420 tectetagaa aggeaaacag tagatttatt teettetttt etagaaacat cattataact 480 aacaatatat aattggaata atggctggtt gggatatttt tggttggtgt atgttatcac 540 cctgttcacg tttttcggat acttagtttt attcaatgtg gtaaacattg aatgtttca 600 gcttaagatc tattttttt tttctagaag aaattgcgtc ctttactaac tttattttac 660 tgtacagtca gagatgtgtt ggcttccctt ggtctgtgga acaaacatgg taaactactt 720 ttcttgggtt tggataatgc cggtaagacc acattgctac atatgttaaa gaacgataga 780 ttggcaacct tacaaccaac atggcatcca acttctgaag aactggctat tggtaacatt 840 aagtttacaa ctttcgattt gggtggtcat attcaagctc gtcgtttatg gaaggattat 900 ttcccagaag ttaatggtat cgtcttttta gtcgatgctg ctgaccctga aagatttgat 960 gaagcacgtg tcgaattaga tgctttattc aacattgccg aattgaagga cgttcctttt 1020 gtaattettg gtaacaagat cgatgeteca aacgeegttt etgaagegga getaegttet 1080 gctttaggat tattgaatac cactggctct caaagaattg aaggtcaaag accagttgaa 1140 gttttcatgt gttccgttgt tatgagaaat ggttatttag aggcgttcca atggttatct 1200 caatatattt aa 1212

<210> 282

<211> 190

<212> PRT

<213> Saccharomyces cerevisiae

<400> 282

Met Ala Gly Trp Asp Ile Phe Gly Trp Phe Arg Asp Val Leu Ala Ser

1 5 10 15

Leu Gly Leu Trp Asn Lys His Gly Lys Leu Leu Phe Leu Gly Leu Asp 20 25 30

Asn Ala Gly Lys Thr Thr Leu Leu His Met Leu Lys Asn Asp Arg Leu 35 40 45

Ala Thr Leu Gln Pro Thr Trp His Pro Thr Ser Glu Glu Leu Ala Ile
50 55 60

Gly Asn Ile Lys Phe Thr Thr Phe Asp Leu Gly Gly His Ile Gln Ala 65 70 75 80

Arg Arg Leu Trp Lys Asp Tyr Phe Pro Glu Val Asn Gly Ile Val Phe 85 90 95

Leu Val Asp Ala Ala Asp Pro Glu Arg Phe Asp Glu Ala Arg Val Glu
100 105 110

Leu Asp Ala Leu Phe Asn Ile Ala Glu Leu Lys Asp Val Pro Phe Val 115 120 125

Ile Leu Gly Asn Lys Ile Asp Ala Pro Asn Ala Val Ser Glu Ala Glu 130 135 140

Leu Arg Ser Ala Leu Gly Leu Leu Asn Thr Thr Gly Ser Gln Arg Ile 145 150 155 160

Glu Gly Gln Arg Pro Val Glu Val Phe Met Cys Ser Val Val Met Arg
165 170 175

Asn Gly Tyr Leu Glu Ala Phe Gln Trp Leu Ser Gln Tyr Ile 180 185 190

<210> 283

<211> 1025

<212> DNA

<213> Saccharomyces cerevisiae

<400> 283

ttctttacaa ttcaccttgc attattgaag gagtgctatt cttcgttttg ccaccctttt 60 gctgttttcc ctcatatcac ataatacccg gagaggctct ttcctgcgac agcgcaacat 120

```
CCaaccatcc taaaggtatg ggtgtactga cgatgcgatt atttcattaa gttctgtctt 180
ttttgtataa atgaaaaaag aacggtgaaa tccatagaaa tacagagagc gacgcaaaca 240
gcgcgcagac tctacgggta atagactcac atccacgtga ccagtttcca atcgaacttt 300
ttcactttgc agggaattat tgtttcacta gcaaaggtag cccacttacc actcagctat 360
gcgaaagttt cattgtttga tacatcttga tagtaaccgc aggcttcttt cttagttcat 420
acatacaaac atagccaaag atgtctgcca aagctcaaaa ccctatgcgt gatttgaaga 540
tcgaaaagtt ggtcttaaac atttctgttg gtgaatctgg tgacagattg accagagcct 600
ccaaggtttt agagcaatta tctggtcaaa ctccagttca atccaaggcc agatacactg 660
tcagaacttt cggtatcaga agaaacgaaa aaattgctgt tcacgttacc gtcagaggtc 720
caaaggctga agaaattttg gaaagaggtt tgaaggtcaa ggaataccaa ttgagagaca 780
gaaacttete tgetaceggt aactteggtt teggtattga egaacacatt gaettgggta 840-
tcaagtatga cccatccatc ggtattttcg gtatggattt ctatgtcgtc atgaacagac 900
caggtgctag agtcactaga agaaagagat gtaagggtac cgttggtaac tcccacaaga 960
caactaagga agacaccgtc tcttggttca agcaaaagta cgatgctgat gttttggaca 1020
                                                              1025
```

<210> 284

<211> 174

<212> PRT

<213> Saccharomyces cerevisiae

<400> 284

Met Ser Ala Lys Ala Gln Asn Pro Met Arg Asp Leu Lys Ile Glu Lys

1 10 15

Leu Val Leu Asn Ile Ser Val Gly Glu Ser Gly Asp Arg Leu Thr Arg
20 25 30

Ala Ser Lys Val Leu Glu Gln Leu Ser Gly Gln Thr Pro Val Gln Ser 35 40 45

Lys Ala Arg Tyr Thr Val Arg Thr Phe Gly Ile Arg Arg Asn Glu Lys 50 55 60

Ile Ala Val His Val Thr Val Arg Gly Pro Lys Ala Glu Glu Ile Leu 65 70 75 80

Glu Arg Gly Leu Lys Val Lys Glu Tyr Gln Leu Arg Asp Arg Asn Phe 85 90 95

Ser Ala Thr Gly Asn Phe Gly Phe Gly Ile Asp Glu His Ile Asp Leu 100 105 110

Gly Ile Lys Tyr Asp Pro Ser Ile Gly Ile Phe Gly Met Asp Phe Tyr 115 120 125

Val Val Met Asn Arg Pro Gly Ala Arg Val Thr Arg Arg Lys Arg Cys 130 135 140

Lys Gly Thr Val Gly Asn Ser His Lys Thr Thr Lys Glu Asp Thr Val 145 150 155 160

Ser Trp Phe Lys Gln Lys Tyr Asp Ala Asp Val Leu Asp Lys 165 170

```
<212> DNA
<213> Candida albicans
<400> 285
agagaacgta ttgaagagga gaaaagggag aaaagaggtc aattggaaga acaacatcgt 60
tctgcatcaa atgcttctat ggcttcttta ttgtcagctg cttcaactac agcagcaact 120
aaaaatttga gtgtggctgg cacaaatcct tctcatacca ctgaaagaat gtttttaaat 180
ttacctttta acaattccct gttcaatgcc ccaccagtag aaattaattt taatgatctt 240
gaagttttgg aattgtacac tcaattagta ttataccgag atgatattac caaatctact 300
tttgaattag ctatatcacc agcaaatttg aatatttctc aacggaaaat catatcaatt 360
ttatgtaatt atttgaattt attagaattg tttgataatg ggttgataat aattagaaga 420
aaaccaggat acattgctca gtgtataact caacaatcta ttattcctaa ttctcaacag 480
gtgtctgggc caactcaccc gcaacaacat caacagaatc aacttcaaca acagcaacag 540
caacaacatc aacatcaaca tccttcacat tcatcatcga tgatgaacct tcatcaattg 600
ggtggtacat tagctgttcc agcgcaccct gaattattaa gatcccaatc gcaatcagca 660
ttaccgttgc caagattgag acagcaaacc tctacaccaa ttcaacaaaa tcaacaagtt 720
cagcaccaaa atcaaccacc acaacaacaa cagcaacagc atgttcaacc acaatataat 780
tattacaatc agcaatctat tcaaagccaa ccacattctg cgagacctta ttctcaatca 840
tataatattt atcaacaaca acagcaacag cagcaacaac aagctcaaca acaagctcaa 900
caacaacaac aacaacaatt acaatatcaa cagggacacc agtcacaagt ttcaacacct 960
acattgaatt cttctagtgc tgctgcatta cttagatcaa gtagcagtag atcatttgtt 1020
gatgtgagat ccacacctcc cacaagtagt tttgctcaac agcaacaaca acaacaacaa 1080
caacagcaac agccgccact aacttcatca atccatgatt ctccaacacc acatcatcat 1140
```

ttaccacttc aacagcagcc accacaacca aatcattacc tatccaatta ccatcagggg 1200

<210> 286 <211> 409

<del><210> 285</del> <211> 1229

<212> PRT

<213> Candida albicans

gttggatctc aaccaaaaac tccattggc

<400> 286

Arg Glu Arg Ile Glu Glu Glu Lys Arg Glu Lys Arg Gly Gln Leu Glu

1 5 10 15

Glı	ı Glı	n His	s Arg		Ala	a Sei	r Asn	Ala 25		Met	Ala	Ser	Leu 30		Ser
Ala	a Ala	a Sei 35		The	Ala	a Ala	Thr		Asn	Leu	Ser	Val 45		Gly	Thr
Asr	Pro 50		His	5 Thr	Thr	Glu 55		Met	Phe	Leu	Asn 60	Leu	Pro	Phe	Asn
Asn	Ser	Ser	Phe	a Asn	Ala	Pro	Pro	Val	Glu	Ile	Asn	Phe	Asn	Asp	Leu
65		<del>,</del> -			70					75					80
Glu	Val	. Leu	Glu	Leu 85		Thr	Gln	Leu	Val 90	Leu	Tyr	Arg	Asp	Asp 95	Ile
Thr	Lys	Ser	Thr 100		Glu	Leu	Ala	Ile 105	Ser	Pro	Ala	Asn	Leu 110	Asn	Ile
Ser	Gln	Arg	Lys	Ile	Ile	Ser	Ile 120	Leu	Cys	Asn	Tyr	Leu 125	Asn	Leu	Leu
Glu	Leu 130	Phe	Asp	Asn	Gly	Leu 135	Ile	Ile	Ile	Arg	Arg 140	Lys	Pro	Gly	Tyr
Ile 145	Ala	Gln	Cys	Ile	Thr 150	Gln	Gln	Ser	Ile	Ile 155	Pro	Asn	Ser	Gln	Gln 160
Val	Ser	Gly	Pro	Thr 165	His	Pro	Gln	Gln	His 170	Gln	Gln	Asn	Gln	Leu 175	Gln
Gln	Gln	Gln	Gln 180	Gln	Gln	His	Gln	His 185	Gln	His	Pro	Ser	His 190	Ser	Ser
Ser	Met	Met 195	Asn	Leu	His	Gln	Leu 200	Gly	Gly	Thr	Leu	Ala 205	Val	Pro	Ala
His	Pro 210	Glu	Leu	Leu	Arg	Ser 215	Gln	Ser	Gln	Ser	Ala 220	Leu	Pro	Leu	Pro
Arg 225	Leu	Arg	Gln	Gln	Thr 230	Ser	Thr	Pro	Ile	Gln 235	Gln	Asn	Gln	Gln	Val 240
Gln	His	Gln	Asn	Gln 245	Pro	Pro	Gln	Gln	Gln 250	Gln	Gln	Gln	His	Val 255	Gln

260

Pro Gln Tyr Asn Tyr Tyr Asn Gln Gln Ser Ile Gln Ser Gln Pro His.

265

Ser Ala Arg Pro Tyr Ser Gln Ser Tyr Asn Ile Tyr Gln Gln Gln Gln 275 280 285

Gln Gln Leu Gln Tyr Gln Gln Gly His Gln Ser Gln Val Ser Thr Pro 305 310 315 320

Thr Leu Asn Ser Ser Ser Ala Ala Ala Leu Leu Arg Ser Ser Ser Ser

<del>325 330 335 335</del>

Arg Ser Phe Val Asp Val Arg Ser Thr Pro Pro Thr Ser Ser Phe Ala 340 345 350

Ser Ser Ile His Asp Ser Pro Thr Pro His His Leu Pro Leu Gln 370 375 380

Gln Gln Pro Pro Gln Pro Asn His Tyr Leu Ser Asn Tyr His Gln Gly 385 390 395 400

Val Gly Ser Gln Pro Lys Thr Pro Leu 405

<210> 287

<211> 2153

<212> DNA

<213> Candida albicans

<400> 287

```
gattgacaaa taataggaat ttacaatcct tgtggataaa ccaaccatct attgctcctt 780
ctaatgttgt tccttgggtg gaacagcaac aacaacaac ccttgatctg ttggaaaata 840
atactaaaac agactccagt aatgatgctt ctgctactaa taataataat gttaatgtta 900
atgttaatgc caatgccaat gttaatgcca atgccaatat ccatgcccaa acccatgtca 960
acacaaatgt taatgcaaac acaacagcaa ctagtattaa tgcttccacg attttaaata 1020
ctacaccaag tattaatgac actaatgata atgccaaaaa aattaatgtt tccatgatta 1080
gtaataacaa taacaataac agtaacaata ataacaacaa cactaacaat agtagtacag 1140
gaagttetaa cattgeaaac atgetteett etgtttetaa tgeaacgaca atgaataaca 1200
gtaatagtat caatagcacc acaaacaata ctacaattaa cgaagctgat gatgacgagt 1260
taattcctac tgcgattgtg attaaaaata ttccatttgc cattaaaaag gaacaattgt 1320
tagatgtgat gacaaaatta aacttgccat taccatatgc ctttaattat cattttgata 1380
atggtgtgtt tegtggatta geatttgeea attteaeate tacagatgag acateageag 1440
tagtaaacca attgaatgga agagaaatag gtggaagaaa attacgtgtt gaatacaaaa 1500
aaatgcttcc ggcacaagaa agagaacgta ttgaaagaga gaaaagggag aaaagaggtc 1560
aattggaaga acaacatcgt tctgcatcaa atgcttcttt ggcttcttta ttgtcagctg 1620
cttcaactac agcagcaact aaaaatttga gtgtggctgg cacaaatcct tctcatacca 1680
ctgaaagaat gtttttaaat ttacctttta acaattccct gttcaatgcc ccaccagtag 1740
aaattaattt taatgatett gaagttitgg aattgtacae teaattagta ttatacegag 1800
atgatattac caaatctact tttgaattag ctatatcacc cagcatattt gaatatttct 1860
caacggaaaa tcatatcaat ttatgtaatt atttgaattt attagaattg tttgataatg 1920
ggttgctact aattagaaaa aaaccaggat ccattgctca gtgtataact caaaaatcta 1980
ttattcctaa ttctcaacag gtgtctgggc caactcaccc gcaccaccat caaaagaatc 2040
aacttcacca acagcaacag caacacacat caacatcaac atccttaaca ttcatcatcg 2100
atgatgaacc ttcatcaatt gggtggtaca ttagctgttc cagcgcacca tga
                                                                  2153
```

```
<210> 288
```

<211> 550

<212> PRT

<213> Candida albicans

<400> 288

Met Asp Phe Arg Asn Leu Ser Thr Thr Pro Asn Gln Met Gly Thr Val 1 5 10 15

Met Gln Arg Arg Pro Ser Leu Ser Ser Leu Ser Ser Ala Ser Gly Tyr
20 25 30

Ser Ser Ser Asn Tyr Gly Gly Asn Pro Thr Pro Asn Pro Asn Asn Ser

35
40
45

Asn Thr Asn Asn Asn Ser Ser Gly Asn Ser Asn Asn Thr His Gly 50 55 60

Asn Asn Thr Pro Lys Leu Ser Thr Gln Arg Leu Thr Asn Asn Arg Asn 65 70 75 80

Leu Gln Ser Leu Trp Ile Asn Gln Pro Ser Ile Ala Pro Ser Asn Val



. 95

Val Pro Trp Val Glu Gln Gln Gln Gln Gln Thr Leu Asp Ser Leu Glu 100 105 110

Asn Asn Thr Lys Thr Asp Ser Ser Asn Asp Ala Ser Ala Thr Asn Asn 115 120 125

Asn Asn Val Asn Val Asn Val Asn Ala Asn Val Asn Ala Asn 130 135 140

Ala Asn Ile His Ala Cln Thr His Val Asn Thr Asn Val Asn Ala Asn 145 150 155 160

Thr Thr Ala Thr Ser Ile Asn Ala Ser Thr Ile Leu Asn Thr Thr Pro 165 170 175

Ser Ile Asn Asp Thr Asn Asp Asn Ala Lys Lys Ile Asn Val Ser Met 180 185 190

Ile Ser Asn Asn Asn Asn Asn Asn Ser Asn Asn Asn Asn Asn Asn Thr 195 . 200 205

Asn Asn Ser Ser Thr Gly Ser Ser Asn Ile Ala Asn Met Leu Pro Ser 210 215 220

Val Ser Asn Ala Thr Thr Met Asn Asn Ser Asn Ser Ile Asn Ser Thr 225 230 235 240

Thr Asn Asn Thr Thr Ile Asn Glu Ala Asp Asp Asp Glu Leu Ile Pro 245 250 255

Thr Ala Ile Val Ile Lys Asn Ile Pro Phe Ala Ile Lys Lys Glu Gln 260 265 270

Leu Leu Asp Val Met Thr Lys Leu Asn Leu Pro Leu Pro Tyr Ala Phe 275 280 285

Asn Tyr His Phe Asp Asn Gly Val Phe Arg Gly Leu Ala Phe Ala Asn 290 295 300

Phe Thr Ser Thr Asp Glu Thr Ser Ala Val Val Asn Gln Leu Asn Gly 305 310 315 320

Arg Glu Ile Gly Gly Arg Lys Leu Arg Val Glu Tyr Lys Lys Met Leu 325 330 335

Pro Ala Gln Glu Arg Glu Arg Ile Glu Arg Glu Lys Arg Glu Lys Arg

345

350

Gly Gln Leu Glu Glu Gln His Arg Ser Ala Ser Asn Ala Ser Leu Ala 355 360 365

Ser Leu Leu Ser Ala Ala Ser Thr Thr Ala Ala Thr Lys Asn Leu Ser 370 380

Val Ala Gly Thr Asn Pro Ser His Thr Thr Glu Arg Met Phe Leu Asn 385 390 395 400

Leu Pro Phe Asn Asn Ser Ser Phe Asn Ala Pro Pro Val Glu Ile Asn 405 410 415

Phe Asn Asp Leu Glu Val Leu Glu Leu Tyr Thr Gln Leu Val Leu Tyr
420 425 430

Arg Asp Asp Ile Thr Lys Ser Thr Phe Glu Leu Ala Ile Ser Pro Ser 435 440 445

Ile Phe Glu Tyr Phe Ser Thr Glu Asn His Ile Asn Leu Cys Asn Tyr 450 455 460

Leu Asn Leu Leu Glu Leu Phe Asp Asn Gly Leu Leu Leu Ile Arg Lys 465 470 475 480

Lys Pro Gly Ser Ile Ala Gln Cys Ile Thr Gln Lys Ser Ile Ile Pro 485 490 495

Asn Ser Gln Gln Val Ser Gly Pro Thr His Pro His His Gln Lys 500 505 510

Asn Gln Leu His Gln Gln Gln Gln His Thr Ser Thr Ser Thr Ser 515 520 525

Leu Thr Phe Ile Ile Asp Asp Glu Pro Ser Ser Ile Gly Trp Tyr Ile 530 535 540

Ser Cys Ser Ser Ala Pro 545 550

<210> 289

<211> 3254

<212> DNA

<213> Candida albicans

<400> 289						
ataaaaaaag	aaatacaatt	aaaaaaattt	tccttctgtg	aaaaggcaat	ttcgggtcta	60
gtagtaaaca	aagcttaata	atttctccca	ttcaaattta	caacggacga	tgcagaaagt	120
agttttcaaa	ttcagtttct	tttcttttat	atatttttc	ttactttctc	ttcctttctt	180
cctgacagta	tattaataaa	tattctttt	aacttattt	tcaatcaaga	aggtactgaa	240
gatatcaatt	aactctcagt	taaatccata	ttaccagttg	tggaataatc	agaagaaaaa	300
aaaaaaagag	agaaaaatca	cgggaattac	gttctcaaca	gaaaataaca	ataattttt	360
tttattcatt	ccaaggtata	acaagaacgt	taggaataat	ataaaattat	caccaaagct	420
gccatcaacg	tgtgtcgaca	accaatcgac	tcctccctta	actagaacca	tagaacctca	480
acatttgttt	ctatagaaaa	atgaagtttg	aaaaaggtaa	agtgagaatt	ttgcctaaac	540
catcccctac	accaaccaac	ccacaaaccc	cattgccatt	acttccagct	caaactaaac	600
ctgtaaactc	aaaaagaaaa	tcagcagcca	gtacacctgg	aaatgaatca	aagaaatcaa	660
gaaaatcaaa	ttctacagct	tcaacaccca	acagtgctac	accaacatca	gtcggaacac	720
ctccacagaa	aacttccaaa	ccaacaggtc	ataggccagt	gacttcatgt	actttttgtc	780
gtcaacataa	aatcaaatgt	aatgcttcag	ataattatcc	aaacccatgt	gaaagatgta	840
aaaaaatggg	tttgaaatgt	gaaattgacc	ccgaatttag	acctcgcaaa	gggtcacaaa	900
tccaatcatt	gaaactggat	gttgatgaat	tgaaggccaa	gattgaaatg	ttgactaaaa	960
atgaatcttt	gctcacacaa	gctttgaatc	aacacaattt	gaaccacgct	tcgcaacagc	1020
					cgtgcattgt	
					ccaattcctt	
					aattctccat	
					tttattttgg	
					atgaccacac	
					cataaatctc	
					cccaaacttt	
					attaaaacac	
					ttacctaatg	
					ttgtcattac	
					caagtaagtc	
					ttttgtgaac	
					tatttattag	
					tcgctatcta	
					gacggtttat	
					gaaagattaa	
					tatattaaat	
					gtcaagtatg	
					atggtgaatg	
					agtgttttca	
					agtgctagac	
					aaagatttgc ttgtataatt	
					accagaatga	
					gaggcaagaa	
-					atcttgccct	
-					acatcaccta	
					gcaaaactga	
					atgcttgaag	
ctactggttc	aacaagagaa	guutagatt	CLLLACCCEC	guadictica	ccatcacaag	2020

caccaactct tcagcagtat ccaatgcagc aagatcaaca acagcaagaa ccatcacaac 2880 aacaacaaca aaagcacctg cagcaactgc agcaatacca acaacagcag caatcgaatc 2940 aacagcaacc acatcttcaa catcaaagac agtttcaaca atcaccacca ccgcaatttt 3000 caatgatttc ttcaacgccg ccccttcagc aaccaccttt tattttggca aactcaccgt 3060 taccacaaca ttatttgcca aagattgatg aaatgaatat gtcaccagaa gtaaaacaag 3120 aaaactctgt tgctccattt gcatccaaa tcaccaactt ttttgatcaa caaacaagtg 3180 gatggtttaa taatgataac caagatgacg actttttggg ttggtttgat gttaatatga 3240 tgcaagagaa ataa 3254

<210> 290

<211> 917

<212> PRT

<213> Candida albicans

<400> 290

Met Lys Phe Glu Lys Gly Lys Val Arg Ile Leu Pro Lys Pro Ser Pro 1 5 10 15

Thr Pro Thr Asn Pro Gln Thr Pro Leu Pro Leu Pro Ala Gln Thr
20 25 30

Lys Pro Val Asn Ser Lys Arg Lys Ser Ala Ala Ser Thr Pro Gly Asn 35 40 45

Glu Ser Lys Lys Ser Arg Lys Ser Asn Ser Thr Ala Ser Thr Pro Asn 50 55 60

Ser Ala Thr Pro Thr Ser Val Gly Thr Pro Pro Gln Lys Thr Ser Lys
65 70 75 80

Pro Thr Gly His Arg Pro Val Thr Ser Cys Thr Phe Cys Arg Gln His
85 90 95

Lys Ile Lys Cys Asn Ala Ser Asp Asn Tyr Pro Asn Pro Cys Glu Arg 100 105 110

Cys Lys Lys Met Gly Leu Lys Cys Glu Ile Asp Pro Glu Phe Arg Pro 115 120 125

Arg Lys Gly Ser Gln Ile Gln Ser Leu Lys Ser Asp Val Asp Glu Leu 130 135 140

Lys Ala Lys Ile Glu Met Leu Thr Lys Asn Glu Ser Leu Leu Thr Gln 145 150 155 160

Ala Leu Asn Gln His Asn Leu Asn His Ala Ser Gln Gln Gln Ser-165 170 175



Ser Gly Ser Gln Ser Gln Gln Gln His Pro Pro Asn Pro Gln Arg Ala 180 185 190

Leu Ser Tyr Thr Ser Ala Asn Ser Ser Pro Gln Val Ala Phe Ser Asn 195 200 205

Ala Ser Pro Ile Pro Ser Val Thr Ser Ile Gln Gln Asn Ala Pro Leu 210 215 220

Thr His Glu Asn Ser Asp Asn Ser Pro Tyr Ala Leu Asn Thr Pro Glu

Asn Ile Glu Glu Leu Gln Pro Ile Ser Glu Phe Ile Leu Gly Asp Val 245 250 255

Thr Leu Pro Leu Asn Arg Ala Asn Glu Leu His Asp Lys Phe Met Thr 260 265 270

Thr His Leu Pro Phe Leu Pro Ile Ile Ile Ser Arg Ser Ala Thr Glu 275 280 285

Leu Tyr His Lys Ser Gln Leu Leu Phe Trp Ala Val Ile Leu Thr Ala 290 295 300

Ser Leu Ser Glu Pro Glu Pro Lys Leu Tyr Met Ser Leu Ala Ser Leu 305 310 315 320

Ile Lys Gln Leu Ala Ile Glu Thr Cys Trp Ile Lys Thr Pro Arg Ser 325 330 335

Thr His Val Ile Gln Ala Leu Ile Ile Leu Ser Ile Trp Pro Leu Pro 340 345 350

Asn Glu Lys Val Leu Asp Asp Cys Ser Tyr Arg Phe Val Gly Leu Ala 355 360 365

Lys Asn Leu Ser Leu Gln Leu Gly Leu His Arg Gly Gly Glu Phe Ile 370 375 380

Gln Glu Phe Ser Arg Asn Gln Val Ser Leu Gly Pro Asp Ala Glu Arg 385 390 395 400

Trp Arg Thr Arg Ser Trp Leu Ala Val Phe Phe Cys Glu Gln Phe Trp 405 410 415

Ser Ser Leu Leu Gly Leu Pro Pro Ser Ile Asn Thr Thr Asp Tyr Leu 420 425 430



				405					100						
Gly	Ser	Leu	Ser.	Leu	Leu	Asp	Arg	Glu	Leu	Glu	Arg	Leu	Arg	Phe	Lys
Ile 465	Ser	Val	Thr	Arg	Pro 470	Asp	Gly	Leu	Leu	Glu 475	Pro	Ser	Asn	Arg	Ala 480
Leu	Ile 450	Ser	Leu	Ser	Ile	Phe 455	Gln	Сув	Lys	Leu	Val 460	Asn	Ile	Met	Gly
Leu	Glu	Asn 435	Ala	Arg	Val	Asp	Lys 440	Ser	Leu	Pro	Lys	Asn 445	Phe	Arg	Cys

485 490 495

Leu Gln Phe Glu Glu Gly Gly Pro Ile Glu Val Tyr Tyr Leu Tyr Ile
500 505 510

Lys Leu Met Ile Cys Cys Phe Ala Phe Leu Pro Gly Thr Pro Ile Glu 515 520 525

Asp Gln Val Lys Tyr Val Ser Phe Ala Tyr Leu Ser Ala Thr Arg Ile 530 535 540

Val Thr Ile Val Ser Lys Met Val Asn Asp Ile Ser Leu Ile Glu Leu 545 550 555 560

Pro Ile Tyr Ile Arg Gln Ala Val Thr Tyr Ser Val Phe Met Leu Phe 565 570 575

Lys Leu His Leu Ser Arg Tyr Leu Ile Asp Lys Tyr Val Asp Ser Ala 580 585 590

Arg Gln Ser Ile Val Thr Val His Arg Leu Phe Arg Asn Thr Leu Ser 595 600 605

Ser Trp Lys Asp Leu Gln Asn Asp Ile Ser Arg Thr Ala Lys Val Leu 610 615 620

Glu Asn Leu Asn Met Val Leu Tyr Asn Tyr Pro Glu Ile Phe Leu Asn 625 630 635 640

Asp Ser Glu Asn Glu Asp Ser Ser Ile Ile Thr Arg Met Arg Ser His 645 650 655

Leu Thr Ala Ser Leu Phe Tyr Asp Leu Val Trp Cys Val His Glu Ala 660 665 670

Arg Arg Ser Val Leu Asp Lys Gly Lys Arg Gln Ala Gln Pro Asn 675 680 685



Lys Lys Ile Leu Pro Leu Pro Phe Tyr Asn Gln Ile Thr Lys Asp Asp 690 695 700

Phe Lys Thr Ile Thr Thr Ser Pro Asn Gly Thr Thr Ile Thr Thr 705 710 715 720

Leu Val Pro Thr Asp Gln Ala Met Asn Gln Ala Lys Ser Lys Ser Phe 725 730 735

Asp Ser Ser Lys Pro Leu Glu Ile Asn Gly Ile Pro Leu Pro Met Leu

Glu Ala Thr Gly Ser Thr Arg Glu Val Leu Asp Ser Leu Pro Ser Gln 755 760 765

Ser Leu Pro Ser Gln Ala Pro Thr Leu Gln Gln Tyr Pro Met Gln Gln 770 780

Asp Gln Gln Gln Glu Pro Ser Gln Gln Gln Gln Gln Lys His Ser 785 790 795 800

Gln Gln Ser Gln Gln Tyr Gln Gln Gln Gln Gln Ser Asn Gln Gln Gln 805 810 815

Pro His Leu Gln His Gln Arg Gln Phe Gln Gln Ser Pro Pro Gln 820 825 830

Phe Ser Met Ile Ser Ser Thr Pro Pro Leu Gln Gln Pro Pro Phe Ile 835 840 845

Leu Ala Asn Ser Pro Leu Pro Gln Thr Tyr Leu Pro Lys Ile Asp Glu 850 855 860

Met Asn Met Ser Pro Glu Val Lys Gln Glu Asn Ser Val Ala Pro Phe 865 870 875 880

Ala Ser Gln Ile Thr Asn Phe Phe Asp Gln Gln Thr Ser Gly Trp Phe 885 890 895

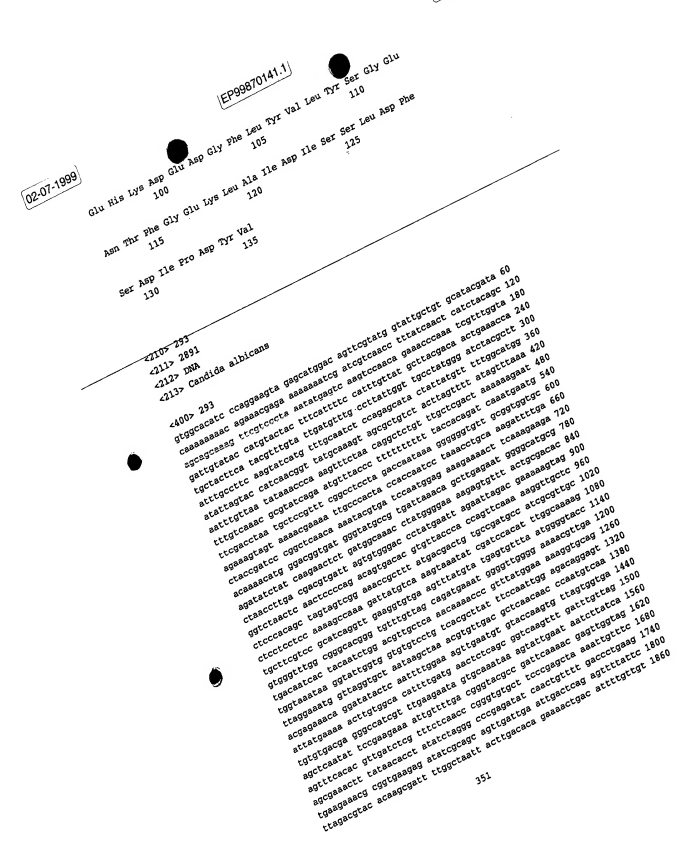
Asn Asn Asp Asn Gln Asp Asp Asp Phe Leu Gly Trp Phe Asp Val Asn 900 905 910

Met Met Gln Glu Lys 915

```
<210> 291
 <211> 908
 <212> DNA
 <213> Candida albicans
 <400> 291
 ttattattac tatgacacac acttactctc ttctatgtct ccgcttacat cacacatcat 60
 acgtttgaaa acacccaccc cctttttctt atctacaaaa aaaaaataac aaactcacta 120
 taactactaa acctttttt ttatttattt attttgtatt ttgttatatt aatcaaacta 180
 ttatcattca tctatatata catccccatc gattcatctg acataatagt atgtatgatc 240
 atacagaggg aaaatcaccc tttttattat taccaaggtc tcaatgatta gtgtggctag 300
 cytclyttac tacttacccc ytyttlytag ataattycca ctcaatcttt aattyaccat 360
cgaccacaaa aataatatgg aaataatact aacagaactt agaaacatca ctcataatga 420
gatcacaatt caaagacgag catccttttg gtatgtatta catgattttt agtcctctga 480
atcttcttcc gatcatcata tactaacatt ttcgtgggtg ttattcttta atagaaaaga 540
gacaagccga ggcagccaga attgctcaga ggttcaagga tagagtacca gtcatctgtg 600
aaaaggttga gaattccgat atccccgaaa ttgataaacg taaatattta gtgccagtgg 660
atttgactgt tggtcaattt gtttacgtta ttagaaaaag aatcaagtta ccaagcgaaa 720
aagccatttt catctttgtc aatgacatat tacccccaac cgctgcatta atcagtacaa 780
tctacgaaga acacaaggac gaagatggtt tcttatacgt tttatactct ggagagaata 840
cttttggcga gaaactagca attgacattt catcattaga tttcagtgat atccctgatt 900
atgtttaa
                                                                   908
<210> 292
<211> 135
<212> PRT
<213> Candida albicans
<400> 292
Met Arg Ser Gln Phe Lys Asp Glu His Pro Phe Glu Lys Arg Gln Ala
Glu Ala Ala Arg Ile Ala Gln Arg Phe Lys Asp Arg Val Pro Val Ile
             20
                                  25
Cys Glu Lys Val Glu Asn Ser Asp Ile Pro Glu Ile Asp Lys Arg Lys
         35
                             40
                                                  45
Tyr Leu Val Pro Val Asp Leu Thr Val Gly Gln Phe Val Tyr Val Ile
     50
                         55
                                              60
Arg Lys Arg Ile Lys Leu Pro Ser Glu Lys Ala Ile Phe Ile Phe Val
 65
                     70
                                          75
                                                              80
```

Asn Asp Ile Leu Pro Pro Thr Ala Ala Leu Ile Ser Thr Ile Tyr Glu

90



Printed: 17-08-2000



<210> 294

taactaatta a

<211> 796

<212> PRT

<213> Candida albicans

<400> 294

Met Phe Thr Leu Phe Phe Phe Thr Thr Asp Gln Met Asn Val Arg Pro

1 5 10 15

Asn Ala Pro Phe Arg Pro Pro Arg Pro Ile Lys Gly Gly Val Ala Val 20 25 30

Val Gln Lys Val Val Lys Arg Lys Leu Pro Thr Thr Thr Asn Pro Lys 35 40 45

Pro Ala Lys Ile Leu Thr Thr Asp Pro Gly Ser Thr Lys Tyr Val Ile
50 55 60

Gln Trp Arg Lys Lys Thr Ser Lys Lys Asn Lys Thr Trp Asp Gly Asp
65 70 75 80

Gly Tyr Ala Val Ile Lys Gln Leu Glu Asn Gly Ala Cys Glu Ile Ser 85 90 95

Ile Lys Asn Ser Asp Gly Lys Pro Met Gly Lys Arg Val Phe Thr Ala 100 105 110

Thr Pro Asn Leu Asp Asp Val Ile Ser Val Gly Pro Tyr Glu Leu Glu

Leu Asp Glu Lys Val Gly Ser Asn Ser Thr Pro Gln Thr Val Thr Arg 130 135 140

125

Val Thr His Gln Phe Lys Lys Val Ala Pro Pro Thr Ala Ser Ser Arg 145 150 155 160

Lys Pro Leu Tyr Asp Asp Cys Ala Asp Ala Ile Ala Leu Pro Pro Pro 165 170 175

Pro Lys Ala Lys Asp Tyr Val Lys Val Asn Ile Asp Pro His Leu Ala 180 185 190

Lys Val Leu Arg Pro His Gln Val Glu Gly Val Lys Phe Met Tyr Glu 195 200 205

Cys Leu Met Gly Tyr Arg Gly Phe Gly Gly His Gly Cys Leu Leu Ala 210 215 220

Asp Glu Met Gly Leu Gly Lys Thr Leu Met Thr Ile Thr Thr Ile Trp 225 230 235 240

Thr Leu Leu Lys Gln Asn Pro Phe Met Glu Lys Gly Ala Val Val Asn 245 250 255

Lys Val Leu Val Val Cys Pro Val Thr Leu Ile Ser Asn Trp Arg Gln 260 265 270

Glu Phe Arg Lys Trp Leu Gly Ala Asn Lys Leu Asn Val Leu Thr Leu 275 280 285

Asn Asn Pro Met Ser Asn Glu Lys Gln Asp Ile Leu Asn Phe Gly Lys

Leu Asn Val Tyr Gln Val Leu Val Val Asn Tyr Glu Lys Leu Val Ala 305 310 315 320

His Phe Asp Glu Leu Ser Ala Val Lys Phe Asp Leu Leu Val Cys Asp 325 330 335

Glu Gly His Arg Leu Lys Asn Ser Ala Asn Lys Val Leu Asn Asn Leu 340 345 350

Ile Lys Leu Asn Ile Pro Lys Lys Ile Val Leu Thr Gly Thr Pro Ile 355 360 365

Gln Asn Glu Leu Val Glu Phe His Thr Leu Ile Ser Phe Leu Asn Pro

SEQL

370

375

380

Gly Val Leu Pro Glu Leu Lys Leu Phe Gln Arg Asn Phe Ile Thr Pro 385 390 395 400

Ile Ser Arg Ala Arg Asp Ile Asn Cys Phe Asp Pro Glu Val Lys Lys
405 410 415

Arg Gly Glu Glu Ile Ser Gln Gln Leu Ile Glu Leu Thr Gln Ser Phe
420 425 430

Ile Leu Arg Arg Thr Gln Ala Ile Leu Ala Asn Tyr Leu Thr Gln Lys
435
440
445

Thr Asp Ile Leu Leu Phe Val Pro Pro Thr Ser Leu Gln Leu Lys Leu 450 455 460

Phe Asp Tyr Ile Thr Asn Leu Lys Lys Phe Asn Gln Phe Glu Ala Phe 465 470 475 480

Thr Met Ile Asn Leu Phe Lys Lys Ile Cys Asn Ser Pro Ser Leu Leu 485 490 495

Ala Asp Asp Glu Leu Phe Lys Lys Ile Val Glu Glu Lys Phe Asn Leu
500 505 510

Gly Met Ala Ser Gly Lys Ile Asn Ile Leu Val Pro Leu Leu Glu 515 520 525

Ile Ala Ser Leu Gly Glu Lys Ile Val Leu Ile Ser Asn Tyr Thr Lys 530 535 540

Thr Leu Asp Leu Leu Glu Gln Val Leu Arg Lys Val Ser Leu Thr Phe 545 550 555 560

Ser Arg Leu Asp Gly Ser Thr Pro Asn Asn Val Arg Ser Lys Leu Val 565 570 575

Asn Gln Phe Asn Thr Asn Pro Asp Ile Asn Val Phe Leu Leu Ser Ser
580 585 590

Lys Ser Gly Gly Met Gly Ile Asn Leu Val Gly Ala Ser Arg Leu Ile 595 600 605

Leu Phe Asp Asn Asp Trp Asn Pro Ala Thr Asp Leu Gln Ser Met Ser 610 615 620

Arg Ile His Arg Asp Gly Gln Leu Lys Pro Cys Phe Ile Tyr Arg Leu



635

640

Phe Thr Thr Gly Cys Ile Asp Glu Lys Ile Phe Gln Arg Gln Leu Val 645 650 655

Lys Asn Lys Leu Ser Ser Lys Phe Leu Asp Asn Asp Ala Thr Ser Lys 660 665 670

Ser Asp Val Phe Asp Asn Asp Asp Leu Lys Asn Ile Phe Glu Ile Asp 675 680 685

Thr Ser Thr Ile Ser Asn Thr His Asp Leu Leu Glu Cys Val Cys Glu
690 695 700

Gly Asp Gly Ser Met Leu Ser Gln Pro Thr Ile Glu Glu Ser Glu Pro 705 710 715 720

Pro Pro Lys Gln Ala Trp Val Thr Ala Leu Glu Leu Lys Lys Lys Ile 725 730 735

Asp Asp Gly Glu Ala Leu Lys Arg Thr Ala Val Lys Phe Ala Leu Asn 740 745 750

Asp Tyr Arg His Tyr Asn Pro Glu Val Asn Arg Asn Leu Asp Phe Asp 755 760 765

Ser Ala Leu His Arg Ile Ala Asn Asn Ser Ser Tyr Glu Asn Lys Gln 770 780

Leu Pro Ile Thr Phe Ile Met Ser Arg Val Thr Asn 785 790 795

<210> 295

<211> 2643

<212> DNA

<213> Candida albicans

<400> 295



```
caattgaata cttcaacgag atgactttac caattcagga tttagaacct gattattata 540
tttccgtcaa ttatcctacc accgataatg gatcaccaac cccacaagct gaaaaatcat 600
tgaaaacatt aattgattta ttatacgata aagggtttgc cgcccaaatt agacctggtg 660
atttagacca tttgttagtc tttgttaaat tgtcttcata caagttttct gaagaagctg 720
aaaaagattt aattaaaaat tatgaatttg gtgtcacggg taaagatgac gtgttagctt 780
ctaaacttag aattatttat caatacttaa cttatccaca atcagttggt ggatgtggta 840
ttactcctaa ttctggggat tggaaatttg tcaccagtat tgttccaatt actaatgcct 900
ttaatgaaac cactttagtt gaagatttaa aaattaatgt tactcaacca aatttatcaa 960
ttgccactat caaaaagaca tatggagttg aagttgctct ttattttgaa tatataaaac 1020
attacacttt ttggttatta ttgctttcta ttattggtct tgtatctcat tttagaaaag 1080
ataaacgatt cctgttaact tttgccttta tcaatttgct ttgggggggtt ttattccttg 1140
catcatggca tagaagagaa caacatttgg ttaatgtatg gggtgtteaa aatagtcatt 1200
taattgaaga acataattcc gaattggcta aagtcaatga aagatatgaa gaaaaatcaa 1260
cttatttcca tgcaaataat accaatggat tcagattttt aaaacaattg gcatttatcc 1320
ccattgcctt ggtgtttgtt ggtgttttga ttagttatca attgagttgt ttctgtattg 1380
aaatcttttt aaccgatatt tatgatggcc ccgggaaatc tttattgact ttattaccaa 1440
cggttttaat cagtgtattt gtgccaattt tgaccattgt ttataatgct gtcacggata 1500
ttattattaa atgggaaaat catgataacc aatatagcaa aaataattct attcttgtta 1560
aaacctttgt gttgaattte ttgactggtt atgttecatt aatcatcact tcattcatat 1620
atttaccatt tgctcatttg gtgcaacctc atttaggtga tattaaaacc actattgcca 1680
catatgctgg tgaaaataga ttctacacca aatacttgtt gaaattaaag agtcaagaag 1740
aatttaaaat caatcaaggt agattagatg ctcaattctt ttatttcatt gtcacaaatc 1800
aagttataca attggtattg aaatatattc teecattggg tttaagattt gtatttaatt 1860
ttattgaaac gaaaattcag aagaaacctc aattacaaac taaagatgat aaccctgatg 1920
aatctatttg gttacataat gtcagattat cgttgaaact tcctgaatat aatgttgatg 1980
atgattttag aggattagtt ttacaatttg gatatttgat aatgtttggt ccagtttggc 2040
cattggcacc attggtttgt attattttca atttaatttt tttcaagttg gataatttta 2100
aattattgaa tggtaaatat ttcaaaccac cagttccaag aagagttgat tctattcatc 2160
catggaattt agcccttttc ttgttagcat ggattggatc aattatttcc cccgtggtca 2220
cggcatttta ccgtcatggt actgctccac caaaatctat gggtcaattt gcccttgata 2280
aagctagtgt tcatgtttca tcctcagttt tcttggtttt attaatgttt gtttcagaac 2340
atggattttt gattttgagt tatcttttat ttgaattctc ttctttgttc aagagtcaag 2400
ttgaatggga aaatgatttt gttgataatg atattaaatt gagacatgat tattattctg 2460
ggaaagtaaa accaacttat aaagtccact cggatgagtt gtgggagaag tttaccccac 2520
aatcaacttt gaatttcact ggtcctaaac caaccgcaga aactgatgat aaagttgaaa 2580
aaattgcttc taccgaagat gcttatctga cttctgcaga aaaatctact actactgcta 2640
                                                                  2643
ctt
```

```
~~~~ <210> 296
```

```
<400> 296
```

```
Met Thr Leu Pro Ile Gln Asp Leu Glu Pro Asp Tyr Tyr Ile Ser Val
1 5 10 15
```

<sup>&</sup>lt;211> 714

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Candida albicans

Asn Tyr Pro Thr Thr Asp Asn Gly Ser Pro Thr Pro Gln Ala Glu Lys
20 25 30

Ser Leu Lys Thr Leu Ile Asp Leu Leu Tyr Asp Lys Gly Phe Ala Ala 35 40 45

Gln Ile Arg Pro Gly Asp Leu Asp His Leu Leu Val Phe Val Lys Leu
50 55 60

Ser Ser Tyr Lys Phe Ser Glu Glu Ala Glu Lys Asp Leu Ile Lys Asn 65 70 75 80

Tyr Glu Phe Gly Val Thr Gly Lys Asp Asp Val Leu Ala Ser Lys Leu 85 90 95

Arg Ile Ile Tyr Gln Tyr Leu Thr Tyr Pro Gln Ser Val Gly Gly Cys
100 105 110

Gly Ile Thr Pro Asn Ser Gly Asp Trp Lys Phe Val Thr Ser Ile Val 115 120 125

Pro Ile Thr Asn Ala Phe Asn Glu Thr Thr Leu Val Glu Asp Leu Lys 130 135 140

Ile Asn Val Thr Gln Pro Asn Leu Ser Ile Ala Thr Ile Lys Lys Thr 145 150 155 160

Tyr Gly Val Glu Val Ala Leu Tyr Phe Glu Tyr Ile Lys His Tyr Thr 165 170 175

Phe Trp Leu Leu Leu Ser Ile Ile Gly Leu Val Ser His Phe Arg 180 185 190

Lys Asp Lys Arg Phe Ser Leu Thr Phe Ala Phe Ile Asn Leu Leu Trp 195 200 205

Gly Val Leu Phe Leu Ala Ser Trp His Arg Arg Glu Gln His Leu Val 210 215 220

Asn Val Trp Gly Val Gln Asn Ser His Leu Ile Glu Glu His Asn Ser 225 230 235 240

Glu Leu Ala Lys Val Asn Glu Arg Tyr Glu Glu Lys Ser Thr Tyr Phe 245 250 255

His Ala Asn Asn Thr Asn Gly Phe Arg Phe Leu Lys Gln Leu Ala Phe 260 265 270



Ile Pro Ile Ala Leu Val Phe Val Gly Val Leu Ile Ser Tyr Gln Leu 275 280 285

Ser Cys Phe Cys Ile Glu Ile Phe Leu Thr Asp Ile Tyr Asp Gly Pro 290 295 300

Gly Lys Ser Leu Leu Thr Leu Leu Pro Thr Val Leu Ile Ser Val Phe 305 310 315 320

Val Pro Ile Leu Thr Ile Val Tyr Asn Ala Val Thr Asp Ile Ile Ile 325 330 335

Lys Trp Glu Asn His Asp Asn Gln Tyr Ser Lys Asn Asn Ser Ile Leu 340 345 350

Val Lys Thr Phe Val Leu Asn Phe Leu Thr Gly Tyr Val Pro Leu Ile 355 360 365

Ile Thr Ser Phe Ile Tyr Leu Pro Phe Ala His Leu Val Gln Pro His 370 375 380

Leu Gly Asp Ile Lys Thr Thr Ile Ala Thr Tyr Ala Gly Glu Asn Arg 385 390 395 400

Phe Tyr Thr Lys Tyr Leu Leu Lys Leu Lys Ser Gln Glu Glu Phe Lys 405 410 415

Ile Asn Gln Gly Arg Leu Asp Ala Gln Phe Phe Tyr Phe Ile Val Thr
420 425 430

Asn Gln Val Ile Gln Leu Val Leu Lys Tyr Ile Leu Pro Leu Gly Leu 435 440 445

Arg Phe Val Phe Asn Phe Ile Glu Thr Lys Ile Gln Lys Lys Pro Gln 450 455 460

Leu Gln Thr Lys Asp Asp Asn Pro Asp Glu Ser Ile Trp Leu His Asn 465 470 475 480

Val Arg Leu Ser Leu Lys Leu Pro Glu Tyr Asn Val Asp Asp Phe
485 490 495

Arg Gly Leu Val Leu Gln Phe Gly Tyr Leu Ile Met Phe Gly Pro Val 500 505 510

Trp Pro Leu Ala Pro Leu Val Cys Ile Ile Phe Asn Leu Ile Phe Phe 515 520 525

Lys Leu Asp Asn Phe Lys Leu Leu Asn Gly Lys Tyr Phe Lys Pro Pro 530 535 540

Val Pro Arg Arg Val Asp Ser Ile His Pro Trp Asn Leu Ala Leu Phe 545 550 555 560

Leu Leu Ala Trp Ile Gly Ser Ile Ile Ser Pro Val Val Thr Ala Phe 565 570 575

Tyr Arg His Gly Thr Ala Pro Pro Lys Ser Met Gly Gln Phe Ala Leu 580 585 590

Asp Lys Ala Ser Val His Val Ser Ser Ser Val Phe Leu Val Leu Leu 595 600 605

Met Phe Val Ser Glu His Gly Phe Leu Ile Leu Ser Tyr Leu Leu Phe 610 615 620

Glu Phe Ser Ser Leu Phe Lys Ser Gln Val Glu Trp Glu Asn Asp Phe 625 630 635 640

Val Asp Asn Asp Ile Lys Leu Arg His Asp Tyr Tyr Ser Gly Lys Val 645 650 655

Lys Pro Thr Tyr Lys Val His Ser Asp Glu Leu Trp Glu Lys Phe Thr 660 665 670

Pro Gln Ser Thr Leu Asn Phe Thr Gly Pro Lys Pro Thr Ala Glu Thr 675 680 685

Asp Asp Lys Val Glu Lys Ile Ala Ser Thr Glu Asp Ala Tyr Ser Thr 690 695 700

Ser Ala Glu Lys Ser Thr Thr Thr Ala Thr 705 710

<210> 297

<211> 1784

<212> DNA

<213> Candida albicans

<400> 297

cggtaattat gtcacaaaaa caaacaatca acatattaaa tcgttatccc aactttgtca 60 gttttactaa caccttttat tttgtgttat acaaattgca caatcaatta ctataacttt 120 tttttgaaac gtgggctctg tttagtttaa cttcttgtag ttttattatt ccgattgggt 180 tagctcaata actgcattc gtacaataat gttaattcaa ttctaaattc cgatgaaccg 240

```
aacacacaaa aaacatccag ttctggagag atttttcaaa acttctatta taaatagaac 300
cctataagtc cataataatt caattgaagg attatttct tttccctttt ctgattactt 360
tcaccaattt tcttctctcc aaaaaaaaca ccttcttcat ggtttctgtt tctaaattaa 420
tcaataacgg gttgttatta actagtcaaa gtgttttcca agatgttgct actccgcaac 480
aagcttctgt gcaacaatac aatatactca attttcttgg cggtagtgcc ccttatattc 540
aaagaaacgg atatgggatt tctactgata tccctgctgg ttgtgaaatt gctcaaattc 600.
aattgtattc aagacatggt gaaagatacc caagtaaaag taatggtaaa agtttagaag 660
caatttatgc taaatttgaa aactacaaag gtacttttaa aggtgatttg tcattcttaa 720
atgattacac ttattttgtc aaagaccaga gtaactatgc taaggaaact agcccaaaaa 780
attctgaagg aacctatgcc ggtacaacca atgccttgcg tcatggtgct gcgtttagag 840
ccaaatatgg atccttatac aaggaaaact caactttacc aatcttcaca tccaattcta 900
acagagtaca tgaaacttca aagtatttcg ctagagggtt tttaggtgat gattatgaag 960
aaggtaaaac tgtcaagttt aacatcatct ctgaagatgc tgatcttggt gccaatagtt 1020
tgactcctag aagtgcatgt tccaagaaca aagaactgag cagtagtact gccaaaaaat 1080
ataacacaac atatttaaat gctattgctg aaagattagt taaaccaaac ccaggtttga 1140
atttgactac aagtgatgtc aacaatttat tcagttggtg tgcttatgaa atcaacgtca 1200
gaggaagttc accattctgt gatttattca ccaatgaaga attcattaag aactcttatg 1260
gtaacgatct ttccaaatat tattctaatg gtgctggtaa taattacacc agaatcattg 1320
gttcagtgat tttgaattca tccttggaac ttttgaagga cactaagaac tctaatcaag 1380
tatggttatc atttgctcat gatactgatt tggaaatttt ccattctgct ttaggattat 1440
tggaaccagc tgaagattta ccaacatctt acatcccatt ccctaaccca tacgtccatt 1500
cttctattgt tccacaaggt gccagaatat acacagaaaa acttcaatgt ggaaacgatg 1560
cttatgttag atacattatc aacgatgctg tcgtgccaat tccaaaatgt gctactggtc 1620
cagggttctc ttgtaaactt gatgattttg aaaatttcgt taaagaaaga attggagatg 1680
ttgactttgt taaacaatgt ggtgtcaata gtacctaccc atctgagctt actttctact 1740
                                                                  1784
gggattataa aaatgtcact tacagtgctc ctttagaatt gtaa
```

<210> 298

<211> 461

<212> PRT

<213> Candida albicans

<400> 298

Met Val Ser Val Ser Lys Leu Ile Asn Asn Gly Leu Leu Leu Thr Ser

1 5 10 15

Gln Ser Val Phe Gln Asp Val Ala Thr Pro Gln Gln Ala Ser Val Gln
20 25 30

Gln Tyr Asn Ile Leu Asn Phe Leu Gly Gly Ser Ala Pro Tyr Ile Gln 35 40 45

Arg Asn Gly Tyr Gly Ile Ser Thr Asp Ile Pro Ala Gly Cys Glu Ile 50 55 60

Ala Gln Ile Gln Leu Tyr Ser Arg His Gly Glu Arg Tyr Pro Ser Lys
65 70 75 80

Ser Asn Gly Lys Ser Leu Glu Ala Ile Tyr Ala Lys Phe Glu Asn Tyr 90

Lys Gly Thr Phe Lys Gly Asp Leu Ser Phe Leu Asn Asp Tyr 110

Phe Val Lys Asp Gln Ser Asn Tyr 120

Ser Glu Gly Thr Tyr Ala Gly Thr Thr Asn Ala Leu Arg His Gly Ala

Ala Phe Arg Ala Lys Tyr Gly Ser Leu Tyr Lys Glu Asn Ser Thr Leu 145 150 155 160

135

140

Pro Ile Phe Thr Ser Asn Ser Asn Arg Val His Glu Thr Ser Lys Tyr 165 170 175

Phe Ala Arg Gly Phe Leu Gly Asp Asp Tyr Glu Glu Gly Lys Thr Val 180 185 190

Lys Phe Asn Ile Ile Ser Glu Asp Ala Asp Leu Gly Ala Asn Ser Leu 195 200 205

Thr Pro Arg Ser Ala Cys Ser Lys Asn Lys Glu Ser Ser Ser Thr 210 215 220

Ala Lys Lys Tyr Asn Thr Thr Tyr Leu Asn Ala Ile Ala Glu Arg Leu 225 230 235 240

Val Lys Pro Asn Pro Gly Leu Asn Leu Thr Thr Ser Asp Val Asn Asn 245 250 255

Leu Phe Ser Trp Cys Ala Tyr Glu Ile Asn Val Arg Gly Ser Ser Pro 260 265 270

Phe Cys Asp Leu Phe Thr Asn Glu Glu Phe Ile Lys Asn Ser Tyr Gly 275 280 285

Asn Asp Leu Ser Lys Tyr Tyr Ser Asn Gly Ala Gly Asn Asn Tyr Thr 290 295 300

Arg Ile Ile Gly Ser Val Ile Leu Asn Ser Ser Leu Glu Leu Leu Lys 305 310 315 320

Asp Thr Lys Asn Ser Asn Gln Val Trp Leu Ser Phe Ala His Asp Thr 325 330 335

Asp Leu Glu Ile Phe His Ser Ala Leu Gly Leu Leu Glu Pro Ala Glu 340 345 350

Asp Leu Pro Thr Ser Tyr Ile Pro Phe Pro Asn Pro Tyr Val His Ser 355 360 365

Ser Ile Val Pro Gln Gly Ala Arg Ile Tyr Thr Glu Lys Leu Gln Cys 370 375 380

Gly Asn Asp Ala Tyr Val Arg Tyr Ile Ile Asn Asp Ala Val Val Pro 385 390 395 400

Ile Pro Lys Cys Ala Thr Gly Pro Gly Phe Ser Cys Lys Leu Asp Asp 405 410 415

Phe Glu Asn Phe Val Lys Glu Arg Ile Gly Asp Val Asp Phe Val Lys
420 425 430

Gln Cys Gly Val Asn Ser Thr Tyr Pro Ser Glu Leu Thr Phe Tyr Trp
435
440
445

Asp Tyr Lys Asn Val Thr Tyr Ser Ala Pro Leu Glu Leu 450 455 460

<210> 299 <211> 1871 <212> DNA

<213> Candida albicans

## <400> 299

gagttttatg ggattgggtt tttcaattaa agactcttcg tgataatgca ataccaaaac 60 caaaatttat ttaacgatgc aataattatt tcaatgtgag ctatccatga atcagtgaga 120 atctttatta tggaagaaat tggtcaaaat ttgacaatgt ttagtaactt tatatcttgg 180 tgttgggatg gtgccacaaa tggagattgt tgagtgtaca tgaaaaatac gtagttaaat 240 tttgtttctt gttttatta tttagccact tttttatccg attctcata ttacctttgg 300 taaagtgata agattaatat catattagta aaacaccccc aatgatgaat gtttgtattt 360 atagccagac tataaaaatt acggggattt aattcgcgac tcacccacgt tctcacacag 420 tttgtgcttt tttcataagt atgatttgaa ccctaaaata tcaactttca taataaacat 480 aattcttcc agacaactaa atggttggtt tatcacgagt acttaatgct gggtttattc 540 taagtggaca atctgtttc caggatgttg ctgcccaca tcaaactac gggtttattc 540 taatatattg caagtacctt ggtggcagcg gtccatatat tcaaaattca gggtatggga 660 tttctactga tataccagaa aaatgcacta ttgaacaagt tcaaatgat agtagacatg 720 gtgagcgatt tcctagtaaa ggagatgga aatactttaa ttcagtgatg gaagttttca 780 agagatatgg tgaatttcat ggagatttat ctttttaaa tgactatgag tatttcgtta 840 ctaatccaga ttattatgaa aaggagacta ctcctaaaaa ctccaaaggt ccatattttg 900

```
gaactacaaa tttattacya catggagctt attttagaaa aagatatcaa tcactatttg 960
accaaaagga gaagcttgtt gtgtttacta gtaattctgg aaggtgttat caaagtggtg 1020
tctatttcgc tcgaggattt ttaggagatg attactcaga agatacagtt gaatttgttg 1080
ttgttgatga agacaaaaaa atgggtggta attcattgac accaagatac gcttgtaaaa 1140
ctttgaatca agatttacac aaagatttgg tgaatcagta cgataagact tatttggacg 1200
atattttatc tagatggcta gtagacaatc ctggattaga tttaagtgca gatcaggtct 1260
cgtcattatt tctttggtgt gcctttgaga ttaacgttag ggggtattct ccattctgca 1320
atctatttac aaaagatgaa tttatcagaa gtgggtaccg aaacgatgtt ggtaattact 1380
atcaaactgg tccaggtaat aatatgacaa aggtaattgg ctcacctatg gtggaagcgt 1440
cgttgaaaat gcttcaagaa gattcaaaaa tttggttgac atttacccat gatactgata 1500
ttgagatgta tttgacatct ttgggattga ttgttccacc aggggatttg cccgttgatc 1560
gagtaccatt teceaateea tataatgeag cagaattttt eeeteaaggt getagaaett 1620
acactgaaaa attgaaatgt ggtgaaaagc aatatgttag atttattgtg aatgatgcag 1680
tttatccata tccggattgt agtggaggtc ctgggtttac ttgtgaattg aatgatttta 1740
tcaaattagt taaaagtcgt ttacatgatg ttgactataa gcttcaatgt gaagtggacg 1800
gaccagegga attgacattt tattgggatt ataaagacag aaagtataat gegeegttaa 1860
                                                                  1871
tagatcagta a
```

<210> 300

<211> 456

<212> PRT

<213> Candida albicans

<400> 300

Met Val Gly Leu Ser Arg Val Leu Asn Ala Gly Phe Ile Leu Ser Gly
1 5 10 15

Gln Ser Val Phe Gln Asp Val Ala Ala Pro His Gln Ala Ser Ile Glu 20 25 30

Gln Tyr Asn Ile Val Lys Tyr Leu Gly Gly Ser Gly Pro Tyr Ile Gln
35 40 45

Asn Ser Gly Tyr Gly Ile Ser Thr Asp Ile Pro Glu Lys Cys Thr Ile
50 55 60

Glu Gln Val Gln Met Ile Ser Arg His Gly Glu Arg Phe Pro Ser Lys 65 70 75 80

Gly Asp Gly Lys Tyr Phe Asn Ser Val Met Glu Val Phe Lys Arg Tyr 85 90 95

Gly Glu Phe His Gly Asp Leu Ser Phe Leu Asn Asp Tyr Glu Tyr Phe
100 . 105 110

Val Thr Asn Pro Asp Tyr Tyr Glu Lys Glu Thr Thr Pro Lys Asn Ser 115 120 125

Lys	130	Pro	туr	Phe	GTĀ	135	Thr	Asn	Leu	Leu	Arg 140	His	GIĀ	Ala	Tyr
Phe 145	Arg	Lys	Arg	Tyr	Gln 150	Ser	Leu	Phe	Asp	Gln 155	Lys	Glu	Lys	Leu	Val 160
Val	Phe	Thr	Ser	Asn 165	Ser	Gly	Arg	Cys	Туг 170	Gln	Ser	Gly	Val	туr 175	Phe
Ala	Arg	Gly		Leu	Gly	Asp	Asp		Ser	Glu	Asp	Thr	Val	Glu	Phe
			180					185					190	•	
Val	Val	Val 195	Asp	Glu	Asp	Lys	Lys 200	Met	Gly	Gly	Asn	Ser 205	Leu	Thr	Pro
Arg	Tyr 210	Ala	Cys	Lys	Thr	Leu 215	Asn	Gln	Asp	Leu	His 220	Lys	Asp	Leu	Val
Asn 225	Gln	Tyr	Asp	Lys	Thr 230	Tyr	Leu	Asp	Asp	Ile 235	Leu	Ser	Arg	Trp	Leu 240
Val	Asp	Asn	Pro	Gly 245	Leu	Asp	Leu	Ser	Ala 250	Asp	Gln	Val	Ser	Ser 255	Leu
Phe	Leu	Trp	Cys 260	Ala	Phe	Glu	Ile	Asn 265	Val	Arg	Gly	Tyr	Ser 270	Pro	Phe
Cys	Asn	Leu 275	Phe	Thr	Lys	Asp	Glu 280	Phe	Ile	Arg	Ser	Gly 285	Tyr	Arg	Asn
Asp	Val 290	Gly	Asn	Tyr	Tyr	Gln 295	Thr	Gly	Pro	Gly	Asn 300	Asn	Met	Thr	Lys
Val 305	Ile	Gly	Ser	Pro	Met 310	Val	Glu	Ala	Ser	Leu 315	Lys	Met	Leu	Gln	Glu 320
Asp	Ser 	Lys	Ile	Trp 325	Leu	Thr	Phe	Thr	His 330	Asp	Thr	Asp	Ile	Glu 335	Met
Tyr	Leu	Thr	Ser 340	Leu	Gly	Leu	Ile	Va1 345	Pro	Pro	Gly	Asp	Leu 350	Pro	Val
Asp	Arg	Val 355	Pro	Phe	Pro	Asn	Pro 360	Tyr	Asn	Ala		Glu 365	Phe	Phe	Pro
Gln	Gly 370	Ala	Arg	Thr	Tyr	Thr	Glu	Lys	Leu	Lys	Cys	Gly	Glu	Lys	Gln

```
      Tyr Val Arg Phe Ile Val Asn Asp Ala Val Tyr Pro Tyr Pro Asp Cys

      385
      390
      395
      400

      Ser Gly Gly Pro Gly Phe Thr Cys Glu Leu Asn Asp Phe Ile Lys Leu 405
      410
      415

      Val Lys Ser Arg Leu His Asp Val Asp Tyr Lys Leu Gln Cys Glu Val 420
      425
      430

      Asp Gly Pro Ala Glu Leu Thr Phe Tyr Trp Asp Tyr Lys Asp Arg Lys 435
      440
      445
```

Tyr Asn Ala Pro Leu Ile Asp Gln 450 455

<210> 301 <211> 1888 <212> DNA <213> Candida albicans

<400> 301

tgtagtataa ataagggtat gaaataccaa catcccagaa tatcaacgag atagaagaga 60 ggagtttcaa tatatatett gtgaataata aettegttet aatteaetat acacaactag 120 acgtgtacac gctcaatctc aggtaaagaa agtttatatt ccatcactat ataacaacaa 180 · tcaggctttg caaaaaaaca tttaaaacta atactggtaa tatggaaata taacgcctcg 240 tagttctacg cacgtggcat cctttatcta tttattcaat ttacccctaa tttatgaatt 300 agcttaataa gagcagtcaa attaacacgg ctcaattaat agtacttaat aatatgaagc 360 cgatcaatta accgatcctt tgaataattt gaaaataaaa taaagtaata taaataggta 420 tgcattttcc ctacatttat ttcctctttc tattttaatt tgtttcctaa acagcaacaa 480 caacaattga aattcaaaaa tggtttctgt ttctaaatta ttgaacaatg gattgttatt 540 agctggtcaa agtgtcttcc aagatgttgc tactccacag caagcttctg tgcaacaata 600 taacatcgtc aattctcttg gcggtagtgc cccttatatt caaagaaacg gatatgggat 660 ttctactgat atccctgctg gttgtgaaat tgctcaaatt caattgtatt caagacatgg 720 tgaaagatac ccaagtaaaa gtaatggtaa aagtttagaa gcaatttatg ctaaatttga 780 aaactacaaa ggtactttta aaggtgattt ggctttctta aatgattata cttattttgt 840 tactgataaa aacaattacg aaaaggaaac tagcccaaaa aattctgaag gaacctatgc 900 cggtacaacc aatgcettge gtcacggtge tgcgtttaga gccaaatatg gateettata 960 caaggaaaat tcaacattac cagttttctc ttccaattca ggtagatgtt accaaacttc 1020 aagatatttt gctagaggat ttttaggtga tgactttaaa gaaggtaaaa ctgtcaagtt 1080 taacatcatt tctgaagatg ctgatgttgg tgccaatagt ttgactccaa gaagtgcatg 1140 ttccaagaac aaagaacgga gcagtagtac tgccaaaaaa tataacacaa catatttaaa 1200 tgctatcact gaaagattag ttaaaccaaa cccaggtttg aatttgacta caagtgatgt 1260 caacaattta ttcagttggt gtgcttatga aatcaacgtc agaggaagtt caccattctg 1320 tgatttattc accaatgaag agtttatcaa atattcttat ggtaatgacc tttccaacta 1380 ttattctaat ggtgctggta acaattacac cagaatcatt ggttcagtga ttttaaattc 1440

```
ttetttagaa ettttaaaag acactaaaaa etetaateaa gtatggttat eatttgetea 1500 tgatactgat ttagaaattt teeattetge tttaggatta ttggaaceag etgaagattt 1560 aceaacatet tacateecat teectaacee atacgteeat teetetattg tteeacaagg 1620 tgeeagaata tacacagaaa aactteaatg tggaaacgat gettatgtta gatacattat 1680 caacgatget gtegtgeeaa tteeaaaatg tgetaetggt eeagggttet ettgtaaact 1740 tgatgatttt gaaaattteg ttaaagaaag aattggagat gttgaetttg ttaaacaatg 1800 tggtgteaat agtacetaee catetgaget tacttetae tgggattata aaaatgteae 1860 ttacaatget eetttaggtg attttaa 1888
```

```
<210> 302
```

<211> 462

<212> PRT

<213> Candida albicans

<400> 302

Met Val Ser Val Ser Lys Leu Leu Asn Asn Gly Leu Leu Leu Ala Gly

1 5 10 15

Gln Ser Val Phe Gln Asp Val Ala Thr Pro Gln Gln Ala Ser Val Gln
20 25 30

Gln Tyr Asn Ile Val Asn Ser Leu Gly Gly Ser Ala Pro Tyr Ile Gln
35 40 45

Arg Asn Gly Tyr Gly Ile Ser Thr Asp Ile Pro Ala Gly Cys Glu Ile
50 55 60

Ala Gln Ile Gln Leu Tyr Ser Arg His Gly Glu Arg Tyr Pro Ser Lys 65 70 75 80

Ser Asn Gly Lys Ser Leu Glu Ala Ile Tyr Ala Lys Phe Glu Asn Tyr 85 90 95

Lys Gly Thr Phe Lys Gly Asp Leu Ala Phe Leu Asn Asp Tyr Thr Tyr 100 105 110

Phe Val Thr Asp Lys Asn Asn Tyr Glu Lys Glu Thr Ser Pro Lys Asn 115 120 125

Ser Glu Gly Thr Tyr Ala Gly Thr Thr Asn Ala Leu Arg His Gly Ala 130 135 140

Ala Phe Arg Ala Lys Tyr Gly Ser Leu Tyr Lys Glu Asn Ser Thr Leu 145 150 155 160

Pro Val Phe Ser Ser Asn Ser Gly Arg Cys Tyr Gln Thr Ser Arg Tyr
165 170 175

Phe	Ala	Arg	Gly 180	Phe	Leu	Gly	Asp	Asp 185	Phe	Lys	Glu	Gly	Lys 190	Thr	Val	
Lys	Phe	Asn 195	Ile	Ile	Ser	Glu	Asp 200	Ala	Asp	Val	Gly	Ala 205	Asn	Ser	Leu	
Thr	Pro 210	Arg	Ser	Ala	Cys	Ser 215	Lys	Asn	Lys	G1u	Arg 220	Ser	Ser	Ser	Thr	
Ala 225	Lys	Lys	Tyr	Asn	Thr 230	Thr	Tyr	Leu	Asn	Ala 235		Thr	Glu	Arg	Leu 240	
	Lys	Pro	Asn	Pro 245		Leu	Asn	Leu	Thr 250	Thr	Ser	Asp	Val	Asn 255	Asn	
Leu	Phe	Ser	Trp 260	Cys	Ala	Tyr	Glu	Ile 265	Asn	Val	Arg	Gly	Ser 270	Ser	Pro	
Phe	Çys	Asp 275	Leu	Phe	Thr	Asn	Glu 280	Glu	Phe	Ile	Lys	Tyr 285	Ser	Tyr	Gly	
Asn	Asp 290		Ser	Asn	Tyr	Tyr 295	Ser	Asn	Gly	Ala	Gly 300	Asn	Asn	Tyr	Thr	
Arg 305	Ile	Ile	Gly	Ser	Val 310	Ile	Leu	Asn	Ser	Ser 315		Glu	Leu	Leu	120 320	
Asp	Thr	Lys	Asn	Ser 325		Gln	Val	Trp	Leu 330		Phe	Ala	His	Asp 335	Thr	
y e.v.	T.e.u	Glu	īle	Phe	His	Ser	Ala	Leu	Gly	Leu	Leu	Glu	Pro	Ala	Glu	

Asp Leu Glu Ile Phe His Ser Ala Leu Gly Leu Leu Glu 345 340

Asp Leu Pro Thr Ser Tyr Ile Pro Phe Pro Asn Pro Tyr Val His Ser 360 355

Ser Ile Val Pro Gln Gly Ala Arg Ile Tyr Thr Glu Lys Leu Gln Cys 375

Gly Asn Asp Ala Tyr Val Arg Tyr Ile Ile Asn Asp Ala Val Val Pro 395 390

Ile Pro Lys Cys Ala Thr Gly Pro Gly Phe Ser Cys Lys Leu Asp Asp 410 405

Phe Glu Asn Phe Val Lys Glu Arg Ile Gly Asp Val Asp Phe Val Lys 430 425 420

```
Gln Cys Gly Val Asn Ser Thr Tyr Pro Ser Glu Leu Thr Phe Tyr Trp
435 440 445
```

Asp Tyr Lys Asn Val Thr Tyr Asn Ala Pro Leu Gly Asp Phe 450 455 460

<210> 303

<211> 1886

<212> DNA

<213> Candida albicans

## <400> 303

```
actacttaaa ttggcatatc caaacaaact tgaagtagga gtttccttat ttttattttg 60
tatttatata tttgattgcg attaatgtca taaattttag ttcggtaatt atgtcacaaa 120
aacaaacaat caacatatta aatcgttatc ccaactttgt cagttttact aacacctttt 180
attittgtgtt atacaaattg cacaatcaat tactataact tittittgaa acgtgggctc 240
tgtttagttt aacttcttgt agttttatta ttccgattgg gttagctcaa taactgcatt 300
tegtacaata atgttaatte aattetaaat teegatgaac egaacacaca aaaaacatee 360
agttctggag agatttttca aaacttctat tataaataga accctataag tccataataa 420
ttcaattgaa ggattatttt cttttccctt ttctgattac tttcaccaat tttcttctct 480
ccaaaaaaaa caccttcttc atggtttctg tttctaaatt aatcaataac gggttgttat 540
taactagtca aagtgttttc caagatgttg ctactccgca acaagcttct gtgcaacaat 600
acaatatact caattttctt ggcggtagtg ccccttatat tcaaagaaac ggatatggga 660
tttctactga tatccctgct ggttgtgaaa ttgctcaaat tcaattgtac tcaagacatg 720
gtgaaagatt cccaacagca agtagtggga aagattatga gaaaatttat gctaaattta 780
aaaactacaa tggtacattc aaaggtgatt tgtcattctt aaatgattac acttattttg 840
tcaaagacca gagtaactat gctaaggaaa ctagcccaaa aaattctgaa ggaacctatg 900
ccggtacaac caatgccttg cgtcatggtg ctgcgtttag agccaaatat ggatccttat 960
acaaggaaaa ctcaacttta ccaatcttca catccaattc taacagagta catgaaactt 1020
caaagtattt cgctagaggg tttttaggtg atgattatga agaaggtaaa actgtcaagt 1080
ttaacatcat ctctgaagat gctgatcttg gtgccaatag tttgactcct agaagtgcat 1140
gttccaagaa caaagaactg agcagtagta ctgccaaaaa atataacaca acatatttaa 1200
atgctattgc tgaaagatta gttaaaccaa acccaggttt gaatttgact acaagtgatg 1260
tcaacaattt attcagttgg tgtgcttatg aaatcaacgt cagaggaagt tcaccattct 1320
gtgatttatt caccaatgaa gaattcatta agaactctta tggtaatgat ctttccaaat 1380
attattctaa tggtgctggt aataattaca ccagaatcat tggttcagtg attttgaatt 1440
catccttgga acttttaaaa gacaccgaga actctaatca agtatggtta tcatttgctc 1500
atgatactga tttagaaatt ttccattctg ctttaggatt attggaacca gctgaagatt 1560
taccaacate ttacatecca tteectaace catacgteca ttettetatt gtteeacaag 1620
gtgccagaat atacacagaa aaacttcaat gtggaaacga tgcttatgtt agatacatta 1680
tcaacgatgc tgtcgtgcca attccaaaat gtgctactgg tccagggttc tcttgtaaac 1740
ttgatgattt tgaaaatttc gttaaagaaa gaattggaga tgttgacttt attaaacaat 1800
gtggtgtcaa tagtacctac ccatctgagc ttactttcta ctgggattat aaaaatgtca 1860
cttacaatgc tcctttagaa ttgtaa
                                                                  1886
```

<210> 304

<211> 461

<212> PRT

<213> Candida albicans

<400> 304

Met Val Ser Val Ser Lys Leu Ile Asn Asn Gly Leu Leu Thr Ser

1 5 10 15

Gln Ser Val Phe Gln Asp Val Ala Thr Pro Gln Gln Ala Ser Val Gln
20 25 30

Gln Tyr Asn Ile Leu Asn Phe Leu Gly Gly Ser Ala Pro Tyr Ile Gln 35 40 45

Arg Asn Gly Tyr Gly Ile Ser Thr Asp Ile Pro Ala Gly Cys Glu Ile 50 55 60

Ala Gln Ile Gln Leu Tyr Ser Arg His Gly Glu Arg Phe Pro Thr Ala 65 70 75 80

Ser Ser Gly Lys Asp Tyr Glu Lys Ile Tyr Ala Lys Phe Lys Asn Tyr 85 90 95

Asn Gly Thr Phe Lys Gly Asp Leu Ser Phe Leu Asn Asp Tyr Thr Tyr 100 105 110

Phe Val Lys Asp Gln Ser Asn Tyr Ala Lys Glu Thr Ser Pro Lys Asn 115 120 125

Ser Glu Gly Thr Tyr Ala Gly Thr Thr Asn Ala Leu Arg His Gly Ala 130 135 140

Ala Phe Arg Ala Lys Tyr Gly Ser Leu Tyr Lys Glu Asn Ser Thr Leu 145 150 155 160

Pro Ile Phe Thr Ser Asn Ser Asn Arg Val His Glu Thr Ser Lys Tyr 165 170 175

Phe Ala Arg Gly Phe Leu Gly Asp Asp Tyr Glu Glu Gly Lys Thr Val 180 185 190

Lys Phe Asn Ile Ile Ser Glu Asp Ala Asp Leu Gly Ala Asn Ser Leu 195 200 205

Thr Pro Arg Ser Ala Cys Ser Lys Asn Lys Glu Ser Ser Ser Ser Thr 210 215 220

Ala 225	Lys	Lys	Tyr	Asn	Thr 230	Thr	Tyr	Leu	Asn	Ala 235	Ile	Ala	Glu	Arg	Leu 240
Val	Lys	Pro	Asn	Pro 245	Gly	Leu	Asn	Leu	Thr 250	Thr	Ser	Asp	Val	Asn 255	Asn
Leu	Phe	Ser	Trp 260	Cys	Ala	Tyr	Glu	Ile 265	Asn	Val	Arg	Gly	Ser 270	Ser	Pro
Phe	Суз	Asp 275		Phe	Thr		Glu 280			Ile		Asn 285	Ser	Tyr	Gly
Asn	Asp 290	Leu	Ser	Lys	Tyr	Tyr 295	Ser	Asn	Gly	Ala	Gly 300	Asn	Asn	Tyr	Thr
Arg 305	Ile	Ile	Gly	Ser	Val 310	Ile	Leu	Asn	Ser	Ser 315	Leu	Glu	Leu	Leu	Lys 320
Asp	Thr	Glu	Asn	Ser 325	Asn	Gln	Val	Trp	Leu 330	Ser	Phe	Ala	His	Asp 335	Thr
Asp	Leu	Glu	Ile 340	Phe	His	Ser	Ala	Leu 345	Gly	Leu	Leu	Glu	Pro 350	Ala	Glu
Asp	Leu	Pro 355	Thr	Ser	Tyr	Ile	Pro 360	Phe	Pro	Asn	Pro	туr 365	Val	His	Ser
Ser	Ile 370	Val	Pro	Gln	Gly	Ala 375	Arg	Ile	Tyr	Thr	Glu 380	Lys	Leu	Gln	Cys
Gly 385	Asn	Asp	Ala	Tyr	Val 390	Arg	туr	Ile	Ile	Asn 395	Asp	Ala	Val	Val	Pro 400
Ile	Pro	Lys	Cys	Ala 405	Thr	Gly	Pro	Gly	Phe 410	Ser	Cys	Lys	Leu	Asp 415	Asp
Phe	Glu	Asn	Phe 420	Val	Lys	Glu	Arg	11e 425	Gly	Asp	Val	Asp	Phe 430	Ile	Lys
Gln	Cys	Gly 435	Val	Asn	Ser	Thr	Туг 440	Pro	Ser	Glu	Leu	Thr 445	Phe	Tyr	Trp
Asp	Tyr 450	Lys	Asn	Val	Thr	Tyr 455	Asn	Ala	Pro	Leu	Glu 460	Leu			

<210> 305 <211> 1635

```
<212> DNA
<213> Candida albicans
<400> 305
atatatatat atttatgtat ttttttattg ttgttcagga attttaaaca tgttcatgaa 60
taatgataat ctatgaacaa attaaagaac tctttggttt catttgcaac caatgtgcgt 120
gacttagggc tatagcccta cttttacttg tacgatactg catattttgt tgttgtgcga 180
atagttagcg taataatctt tttttttgtg tgtgtgcggt ttacttactc ttcttctctt 240
cgcacatatt ttattagagc ttacagttgt ttgtatagtg agagtttcac taacacaaag 300
cttcaacaat actaacaaat tttcgcacgc tgtggaagga gaaacttaca ctgtacacta 360
cactacactg tacactatac accaccaaca gaaaaaaaaa attatcaaat tttcaacctt 420
gagagaaaaa aaaaagtgga aaaaaaactt cttcttacat ttagttaatt ttcagacagg 480
cacaaaggaa ttaatcacca tgaaggtatg tgattgaata taacctatat cagtgattat 540
aattagagtc tttatttgga tattgcaata attggataat aaagaaagag cataagagta 600
ggagttttaa acaggataat tggattcaat aagaggaaaa attttttatc gtcgtgatta 660
taacaaatac aaagaaatta agcaatgaag tgatataagc aaatgaagga ctagtttatt 720
aggggtgaca tttttagact acgtaaaagt actttcgatt caaggaaaac caaattttag 780
tatctatcaa caaactacaa atcaatttag ttaacttcaa taatgacaat aattttaatc 840
actgaaaaca ataaacatgc aaacaagcaa aactagtcaa gctttacgaa tcagtcaata 900
ctaacaatac ttttttttgt ttcattttag ttaaacatct catatccagc caacggtact 960
caaaaatcta tggatatcga tgatgacaca aaattacgtg tttctacgga aaaaagaatg 1020
ggtcaagaag ttgaaggtga ctcagttgga gatgaattca aaggttacat cttcaaaatc 1080
actggtggta acgataaaca aggtgtccca atgaaacaag gtgttatgca cccaaccaga 1140
qttagattat tattatctaa aggtcactct tgttacagac caagaagaac tggtgaaaga 1200
aaaagaaaat ccgttagagg ttgtattgtt gctcaagatt tgtcagtttt ggctttgtct 1260
attgttaaac aaggtgacaa tgaaattgaa ggattaactg acaccactgt tccaaaaaga 1320
ttaggtccaa agagagctaa ccacattaga aaattctttg gtttaactaa agaagatgat 1380
gttagagatt tcgttgttag aagagaagtt actaaaggtg acaaaactta caccaaagct 1440
ccaaagattc aaagattagt tactccacaa actttacaaa gaaagagagc tttgaaagct 1500
aaaaaagtca agaatgctca acaacaaaga gatgctgctg ctgaatacgc tcaattgttg 1560
gctaagagat tgcatgaaag aaaagaagaa agagctgaaa ttaaaaagaa gagagctgaa 1620
                                                                  1635
tctttaaaga actaa
<210> 306
<211> 236
<212> PRT
<213> Candida albicans
<400> 306
Met Lys Leu Asn Ile Ser Tyr Pro Ala Asn Gly Thr Gln Lys Ser Met
                                     10
```

20

Asp Ile Asp Asp Asp Thr Lys Leu Arg Val Ser Thr Glu Lys Arg Met 25

Gly Gln Glu Val Glu Gly Asp Ser Val Gly Asp Glu Phe Lys Gly Tyr 35 40 45

Ile Phe Lys Ile Thr Gly Gly Asn Asp Lys Gln Gly Val Pro Met Lys 50 55 60

Gln Gly Val Met His Pro Thr Arg Val Arg Leu Leu Ser Lys Gly 65 70 75 80

His Ser Cys Tyr Arg Pro Arg Arg Thr Gly Glu Arg Lys Arg Lys Ser 85 90 95

Val Arg Gly Cys Ile Val Ala Gln Asp Leu Ser Val Leu Ala Leu Ser 100 105 110

Ile Val Lys Gln Gly Asp Asn Glu Ile Glu Gly Leu Thr Asp Thr Thr
115 120 125

Val Pro Lys Arg Leu Gly Pro Lys Arg Ala Asn His Ile Arg Lys Phe 130 135 140

Phe Gly Leu Thr Lys Glu Asp Asp Val Arg Asp Phe Val Val Arg Arg 145 150 155 160

Glu Val Thr Lys Gly Asp Lys Thr Tyr Thr Lys Ala Pro Lys Ile Gln 165 170 175

Arg Leu Val Thr Pro Gln Thr Leu Gln Arg Lys Arg Ala Leu Lys Ala 180 185 190

Lys Lys Val Lys Asn Ala Gln Gln Gln Arg Asp Ala Ala Ala Glu Tyr
195 200 205

Ala Gln Leu Leu Ala Lys Arg Leu His Glu Arg Lys Glu Glu Arg Ala 210 215 220

Glu Ile Lys Lys Lys Arg Ala Glu Ser Leu Lys Asn 225 230 235

<210> 307

<211> 1520

<212> DNA

<213> Candida albicans

<400> 307

gtgacgagaa cttctgtcat ctcgagtctg ccaactgcct ctaacagcaa caacaataag 60

```
aacaatgata acggaggagg attatcccat acaaacagaa tagttgttgg tgtagttgtt 120
ggggttggtg gttctatatt aattggtttg ttggccgttt tattttactt gagaaagaga 180
aacaaccgtg attatgaagg tggatggact ttctggagaa agaatgagaa attgggaagt 240
gatgagttct tcaatggtga attgggtgtc agagacagaa atattaatca aggatcaaat 300
ttttaaacaa ggcttatttt ggatgagggt ggttttttt ataagtattt tgtagttgaa 360
tttaaaattt tgtaccttaa agtcttttaa tttaatttta ataaaaagtg gtgatttggc 420
aaacttcaag agtatatttg gtgaaaaaaa aaaaaaaatt tggaactgaa cgcgtctaac 480
atcttatacc tctaagcaaa atgtcagagt actctgtgta tcaacagttg aatgaagata 540
caaacgcaac taaatatact tataaattac tacagctacc atcaaagata ctaaatcaac 600
ttgaatccaa gtcaactaac ttgtatataa aatctgatat caattcccta gcattatgca 660
ctgattcaga aactttcaag ttacgacaaa tgaaccattc caatacagtc ttgctattga 720
acaaagaace tgacaacaag ttaattgggt ttcagaaaac cagttatgaa tatgagttga 780
cagaaatcaa aggttcgatc gatacgtccg atatccctat tttcaacgga caaacagcac 840
agcaacctat tgatttgata gcattggaag ataattcgat ttgttcacat caagagtttt 900
tatcgaattg gtatgagttg ggaggttgtg aaattgataa tggagcatat ataatgagtg 960
cagatattat tactgaacta ttatatctat taatcaccaa attgatgagt ttacaagtgc 1020
acgagttttc tccggaagat gtttcatcca tcatcacgcc cccttataat gactcaatgg 1080
taacatcaat catacacaaa ttttgcacta tagaaagtga gaaatatcaa ttgaatgatt 1140
taaaaattac acagtggttt ggcattgttg agatgtcaaa aatcaatcat aaaatgaccg 1200
atatttcaga gttcttattg aattggaaaa ctagtttgcc gtcattctat aaccctccat 1260
tggacatcag tcaattggca ggctattact gctccccaat cgaaaacaaa atattgtatg 1320
tcqacccaga atctttatca gaaaatttga gtcaacgatt caaagaattg tttgaattgg 1380
ataaaagttg gaactatgat gagtttattc cattcattaa aaagtttgtt cctgccggta 1440
aaaaggtcga ctcaattatt ttaaagtatg gcaagaagaa gaaagttggt agagatagat 1500
                                                                  1520
ttatagtctg tcctagataa
```

<210> 308

<211> 339

<212> PRT

<213> Candida albicans

<400> 308

Met Ser Glu Tyr Ser Val Tyr Gln Gln Leu Asn Glu Asp Thr Asn Ala 1 5 10 15

Thr Lys Tyr Thr Tyr Lys Leu Leu Gln Leu Pro Ser Lys Ile Leu Asn 20 25 30

Gln Leu Glu Ser Lys Ser Thr Asn Leu Tyr Ile Lys Ser Asp Ile Asn 35 40 45

Ser Leu Ala Leu Cys Thr Asp Ser Glu Thr Phe Lys Leu Arg Gln Met 50 55 60

Asn His Ser Asn Thr Val Leu Leu Leu Asn Lys Glu Pro Asp Asn Lys
65 70 75 80

I	eu	Ile	Gly	Phe	Gln 85	Lys	Thr	Ser	Tyr	Glu 90	Tyr	Glu	Leu	Thr	Glu 95	Ile
I	ys	Gly	Ser	Ile 100	Asp	Thr	Ser	Asp	Ile 105	Pro	Ile	Phe	Asn	Gly 110	Gln	Thr
A	la	Gln	Gln 115	Pro	Ile	Asp	Leu	Ile 120	Ala	Leu	Glu	Asp	Asn 125	Ser	Ile	Сув
S	er	His 130	Gln	Glu	Phe	Leu	Ser 135	Asn	Trp	Tyr	Glu	Leu 140	Gly	Gly	Суз	Glu
	le .45	Asp	Asn	Gly	Ala	Туг 150	Ile	Met	Ser	Ala	Asp 155	Ile	Ile	Thr	Glu	Leu 160
I	eu	Tyr	Leu	Leu	11e 165	Thr	Lys	Leu	Met	Ser 170	Leu	Gln	Val	His	Glu 175	Phe
S	er	Pro	Glu	Asp 180	Val	Ser	Ser	Ile	Ile 185	Thr	Pro	Pro	Tyr	Asn 190	Asp	Ser
M	iet	Val	Thr 195	Ser	Ile	Ile	His	Lys 200	Phe	Cys	Thr	Ile	Glu 205	Ser	Glu	Lys
1	уr	Gln 210	Leu	Asn	Asp	Leu	Lys 215	Ile	Thr	Gln	Trp	Phe 220	Gly	Ile	Val	Glu
	let 25	Ser	Lys	Ile	Asn	His 230	Lys	Met	Thr	Asp	11e 235	Ser	Glu	Phe	Leu	Leu 240
A	sn	Trp	Lys	Thr	Ser 245	Leu	Pro	Ser	Phe	Tyr 250	Asn	Pro	Pro	Leu	Asp 255	Ile
S	er	Gln	Leu	Ala 260	Gly	Tyr	Tyr	Cys	Ser 265	Pro	Ile	Glu	Asn	Lys 270	Ile	Leu
I	λr	Val	Asp 275	Pro	Glu	Ser	Leu	Ser 280	Glu	Asn	Leu	Ser	Gln 285	Arg	Phe	Lys
G	Slu	Leu 290	Phe	Glu	Leu	Asp	Lys 295	Ser	Trp	Asn	Tyr	Asp 300	Glu	Phe	Ile	Pro
	?he 805	Ile	Lys	Lys	Phe	Val 310	Pro	Ala	Gly	Lys	Lys 315	Val	Asp	Ser	Ile	11e 320
I	Leu	Lys	Tyr	Gly	Lys 325	Lys	Lys	Lys	Val	Gly 330	Arg	Asp	Arg	Phe	11e 335	Val

Cys Pro Arg

<210> 309

<211> 710

<212> DNA

<213> Candida albicans

<400> 309

```
aaagaccaaa aaattattat ttaaaaaaat tttttaccat ccaaaggatt ttattaccc 60 aaagaccaaa aaattattat ttgaaatggg gattaaaatg gggataattt tattttgcct 120 gatgaaaaga ttagtacccg tttgaaagtc ctgattcaaa ttatggtcaa gctaaaagag 180 aaatatttga tttaattact ggtgataaaa aaccaagaat tttccatgtt gatgaatcat 240 taattgatga ttctcaagaa gctgctgatg aatatgaaaa acaattgatt aataattttg 300 ctaaaaaaga ttcggtgaaa ttacctttat ttgatttatt tttattaggt tgtgcaccgg 360 atggtcatat tgcttcatta tttcctaatc atggtgaaca attgagagaa aaattagctt 420 gggttttacc agtatcaaat gctcctctgg gaccagaaaa tagaataact ttatctatcc 480 cagttatatg tcattccgca agagtgacat ttgttgttga agggttaact aaggcaccaa 540 ttattaaaac cattatggaa agaccagaaa aaggtttacc aagttcaatt gttaatgaag 600 gtgctgctgg tagagtgagt tggtttgttg atgatgatgc attgaatgat ttgtttgata 660 taactaaaaa gaaatacaaa tatttatcta tacctgaacc aagtcattaa
```

<210> 310

<211> 131

<212> PRT

<213> Candida albicans

<400> 310

Val Lys Leu Pro Leu Phe Asp Leu Phe Leu Leu Gly Cys Ala Pro Asp

1 5 10 15

Gly His Ile Ala Ser Leu Phe Pro Asn His Gly Glu Gln Leu Arg Glu 20 25 30

Lys Leu Ala Trp Val Leu Pro Val Ser Asn Ala Pro Ser Gly Pro Glu
35 40 45

Asn Arg Ile Thr Leu Ser Ile Pro Val Ile Cys His Ser Ala Arg Val 50 . 55 60

Thr Phe Val Val Glu Gly Leu Thr Lys Ala Pro Ile Ile Lys Thr Ile
65 70 75 80

Met Glu Arg Pro Glu Lys Gly Leu Pro Ser Ser Ile Val Asn Glu Gly 85 90 95

```
Ala Ala Gly Arg Val Ser Trp Phe Val Asp Asp Asp Ala Leu Asn Asp 100 105 110
```

Leu Phe Asp Ile Thr Lys Lys Lys Tyr Lys Tyr Leu Ser Ile Pro Glu 115 120 125

Pro Ser His

<210> 311

```
<211> 1190
<212> DNA
<213> Candida albicans
<400> 311
tgttacaaaa cattctgttg gagagataat tgaattcaaa attaacgtgt tgaatccaac 60
attgaacaat gaatacacgt ccaagaaaaa tttgacatga ttagaatcgc ggtcaattac 120
attcccggaa cgtcttggac tacttggata caacaatgga aaatgaggaa aatgaggaaa 180
atgaggaaaa cgaggaaaac gaggaaaata tttaccgaag agtaattata ttacaagcat 240
aagacaacag ctaaaatttt ttggtcagaa cacaactttg gaagaaagaa aaaaaccgga 360
tttaaaaact acacttcatc atggctggag ttagacaatt aagaataata gcattaacgg 540
cctttgtcct tggtttaatt tttactttac ataaagttgg atccaacgct gcatccttgg 600
ttcatgcaca agcatcagac caacaaccaa acaaacataa caccaaaagt actacatata 660
ccgccactaa tgacgaatca gttgccaatc tcattgattc taaaaatgat cctcaaactg 720
atgacaaaat aaatcaaaaa atatcacaag atcaagatga agccatcaat ggtaataaag 780
acactaataa agacaccacc aaagtcaaac cagataatgg tgaatatgat ccaatatctg 840
atttgataaa aattagatca ttatcaccaa tgacaatttt cagtaaatca tattgtccat 900
attcaaaaaa gattaaacaa ttgttattag aaaaatatga tataacacca gcaccaaatg 960
ttgttgaatt agatcgatat gaatatggag ctgaattaca aagttatttg acagagaaga 1020
gtgggagaag aactgtgcca aacgtattgg ttggtaaatc atttgaaagt aggggtggtt 1080
gtgatgaatt tgaaaaactt cataaagata atgatttgat taaattgtta gttgaatggg 1140
ggtctggtcg tttacaagtt gcaaagaaga ataccccatc aaatgcctaa
                                                        1190
```

```
<210> 312 ·
<211> 229
<212> PRT
```

<213> Candida albicans

<400> 312

Met Ala Gly Val Arg Gln Leu Arg Ile Ile Ala Leu Thr Ala Phe Val 1 5 10 15

Leu Gly Leu Ile Phe Thr Leu His Lys Val Gly Ser Asn Ala Ala Ser 20 25 30

Leu Val His Ala Gln Ala Ser Asp Gln Gln Pro Asn Lys His Asn Thr 35 40 45

Lys Ser Thr Thr Tyr Thr Ala Thr Asn Asp Glu Ser Val Ala Asn Leu 50 55 60

Ile Asp Ser Lys Asn Asp Pro Gln Thr Asp Asp Lys Ile Asn Gln Lys

Ile Ser Gln Asp Gln Asp Glu Ala Ile Asn Gly Asn Lys Asp Thr Asn 85 90 95

Lys Asp Thr Thr Lys Val Lys Pro Asp Asn Gly Glu Tyr Asp Pro Ile 100 105 110

Ser Asp Leu Ile Lys Ile Arg Ser Leu Ser Pro Met Thr Ile Phe Ser 115 120 125

Lys Ser Tyr Cys Pro Tyr Ser Lys Lys Ile Lys Gln Leu Leu Glu 130 135 140

Lys Tyr Asp Ile Thr Pro Ala Pro Asn Val Val Glu Leu Asp Arg Tyr 145 150 155 160

Glu Tyr Gly Ala Glu Leu Gln Ser Tyr Leu Thr Glu Lys Ser Gly Arg 165 170 175

Arg Thr Val Pro Asn Val Leu Val Gly Lys Ser Phe Glu Ser Arg Gly 180 185 190

Gly Cys Asp Glu Phe Glu Lys Leu His Lys Asp Asn Asp Leu Ile Lys 195 200 205

Leu Leu Val Glu Trp Gly Ser Gly Arg Leu Gln Val Ala Lys Lys Asn 210 215 220

Thr Pro Ser Asn Ala 225

<210> 313

<211> 1256

<212> DNA

<213> Candida albicans

```
<400> 313
agtggttgtt caataatggt aagttettgg aaatageeat tgttgettte tggtggttag 60
acttgtagga agtagaactg ttttccaatg aaaagtagtt ttaattagaa aaattttcaa 120
agtgcgtgaa gcccagtctg aatgtgcgag gaagcccagt cagttagtag tgtccttccc 180
tccactgtct gtaatacaaa atttccctta gtgaaaatgc gaaatatatc tgtactggga 240
acccccgga aaaaaaaaaa cctatgctca aaactatatg tactgtacac aatctagggc 300
tatagcccta atattgtaca ggaagaactt taactatggt gcgaagagcg tttccaattt 360
ttttttttc aggtgtagtc tgttctatgg caatactgtt gttagtagag agtgtctcgc 420
actaacagaa catttttttc agaacaggaa aatttttgaa atctaacatc ttttactgaa 480
agecaageat caacacaata atgtcaaccc aatctgttca agtatgtaaa cgaattgaaa 540
taaagagata gagagatgtt ttattatcaa aatacgaaag gaaaggcaat taaaaaagga 600
aatcaaaaag tcccaacctt gcagtagaag aattgaggta tatgaatttg atagatagcc 660
agaacggtgt tacataaatg ggatatagaa caaaactata cgaggagttt gtttcaacga 720
tcattcaata accagaaaac gataatattt tagcgaccat taaatgacac ttgaaggctc 780
actgggccaa tagaatatct ccatatacac ttttgaacta tttactaaca atttactttt 840
gtttctagac ttttggtaaa aagaagactg ccactgccgt tgctcatgtt aaagccggta 900
aaggtttaat taaaattaac ggttccccaa tcaccttggt ccaaccagaa atcttaagat 960
tcaaagttta cgaaccattg actttggttg gtttagataa attccaaggt atcgacatca 1020
gagttaaagt cactggtggt ggtcacgttt ctcaagtcta cgccatcaga caagctattg 1080
ctaaaggttt ggttgcttac caccaaaaat acgttgacga agcttctaag aacgaattaa 1140
agaaaatttt cgcttcttac gataagacct tgttagttgc cgactcaaga agaatggaac 1200
caaagaaatt cggtggtcgt ggtgccagag caagattcca aaaatcttac cgttaa
```

<210> 314

<211> 142

<212> PRT

<213> Candida albicans

<400> 314

Met Ser Thr Gln Ser Val Gln Thr Phe Gly Lys Lys Lys Thr Ala Thr

1 5 10 15

Ala Val Ala His Val Lys Ala Gly Lys Gly Leu Ile Lys Ile Asn Gly
20 25 30

Ser Pro Ile Thr Leu Val Gln Pro Glu Ile Leu Arg Phe Lys Val Tyr
35 40 45

Glu Pro Leu Thr Leu Val Gly Leu Asp Lys Phe Gln Gly Ile Asp Ile 50 55 60

Arg Val Lys Val Thr Gly Gly Gly His Val Ser Gln Val Tyr Ala Ile
65 70 75 80

Arg Gln Ala Ile Ala Lys Gly Leu Val Ala Tyr His Gln Lys Tyr Val

90

95

Asp Glu Ala Ser Lys Asn Glu Leu Lys Lys Ile Phe Ala Ser Tyr Asp 100 105

Lys Thr Leu Leu Val Ala Asp Ser Arg Arg Met Glu Pro Lys Lys Phe 120

Gly Gly Arg Gly Ala Arg Ala Arg Phe Gln Lys Ser Tyr Arg 135

<210> 315

<211> 959

<212> DNA

<213> Candida albicans

<400> 315

gtggtaagat atagaaagct taccactttg acaagtttga aataggatgg gtgaaaattt 60 ggacatettg aatacttaaa attetgaact tgatcaccag atcetttet titacataat 120 tagatatgat ggataggtta gaatcgtctt taaagagaag gtataatatc taactgattt 180 ggcgaggtgt tggaaaagtc actccactgt atatattctc ggagtttaac gtactacagt 240 tcagtggggt gaatacctaa ataggggggt agaatacgaa ctcctacaaa ttttaaggag 300 actatgaccc gaaaagagaa gaaaaattta ttactctaag aactttatat acctccacaa 360 ctcacttttt ctttagtttc attctgcttt ttttttctta cacatcttaa ggtcaaacaa 420 tttaacttat tagcttgtga aaatctcact tcaattcaag ttctctttca attgacatta 480 tagtatttcc caattcaatt atggcttctc atgcttcctg tatattctgt aaaattatca 540 aaggtgaaat teettette aagttaattg aaactgeaaa gaettattee ttettggaca 600 ttcaaccaat tgctgaagcc cacgttttaa ttatccctaa acaccatggg gcaaagttgc 660 acaacattcc agacgactac cttagtgaca ttttaccagt tgtcaaaaaa ttgacaaaag 720 tcttgaaatt ggacgaaaat aatactccag aaggtgaagg ttataacgtt ttacagaaca 780 acggaagaat tgctcatcaa gttgttgatc acgttcactt ccatttgatt cctaaaaaagg 840 atgaggetae aggtttaggt gttggttgge etgetgaage caetgatttt gataaattag 900 gaaaattgca tgagaaatta aaggaagaat tggctaaggt agataatgaa aaattataa 959

<210> 316

<211> 152

<212> PRT

<213> Candida albicans

<400> 316

Met Ala Ser His Ala Ser Cys Ile Phe Cys Lys Ile Ile Lys Gly Glu 1 5 10 15

Ile Pro Ser Phe Lys Leu Ile Glu Thr Ala Lys Thr Tyr Ser Phe Leu 20 25 30

Asp Ile Gln Pro Ile Ala Glu Ala His Val Leu Ile Ile Pro Lys His
35 40 45

His Gly Ala Lys Leu His Asn Ile Pro Asp Asp Tyr Leu Ser Asp Ile
50 55 60

Leu Pro Val Val Lys Lys Leu Thr Lys Val Leu Lys Leu Asp Glu Asn 65 70 75 80

Asn Thr Pro Glu Gly Glu Gly Tyr Asn Val Leu Gln Asn Asn Gly Arg

85 90 95

Ile Ala His Gln Val Val Asp His Val His Phe His Leu Ile Pro Lys
100 105 110

Lys Asp Glu Ala Thr Gly Leu Gly Val Gly Trp Pro Ala Glu Ala Thr
115 120 125

Asp Phe Asp Lys Leu Gly Lys Leu His Glu Lys Leu Lys Glu Glu Leu 130 135 140

Ala Lys Val Asp Asn Glu Lys Leu 145 150

<210> 317

<211> 297

<212> DNA

<213> Candida albicans

<400> 317

<210> 318

<211> 25

<212> PRT

<213> Candida albicans

<400> 318

Met-Arg Asp Lys Trp Arg Lys Lys Arg Val Arg Arg Leu Lys Arg Lys 1 5 10 15

Arg Arg Lys Val Arg Ala Arg Ser Lys
20 25

<210> 319 <211> 1303 <212> DNA

<213> Candida albicans

## <400> 319

tattgtctga tgctatacgg aatgggcgtt acaaatatac aaacttatat ttgaaagtaa 60 attictattat titictictat cgtatgcata ccgattatta tcacaaggac aattgcctat 120 tgttgtgtgg aataaattta aaatccttct tattggtgtc tagactttgc tttttgtggt 180 gattagggct ttagccctat cacgtgaaat actgtatata aaaaattctt tatagcgcga 240 taaaacatat ttttttccg tattaacaaa tatgtgtgaa gttttgtcct ggtgttttct 300 cactgttttt ccttttttt ctggtagtat caattaacgc ttagatccaa tacagttttg 360 qtaacttgta cacgaacaaa atctcaaatt tgttactgtg tgaaccaaca aggaagagaa 420 aaaaaaaccc atacaaaaat ttttcagtat caaggaatta gaagagacgt ttaaatcaac 480 aaagttcaaa tctatcaaca atggtatgtt taatatcgat attatccata gatgtacatg 540 tatcctaatg ggtttcatta tttggaaagt tatgtttatg ggagttctat ttattaagat 600 atgggataag aattaaagta ttggatgagt agtacaagac caacaaagag aaatagcccc 660 ctttccctcc actattcaat atactcaaca acattatcaa gttaaaagtt cagaagatac 720 acgtaaatga aaagttaata ccaagaagaa tacaaattac cagtccatac cgtgtttggg 780 tttagattac tatattttac aagaaacata ttatatgaaa tgatacccaa tccacagcga 840 cttttcagat agccaaataa ctaagcaact caagataaca taggatcatg catcaatcac 900 aaatgaaaca ttaatactaa ctaacttttt ttttatttat taggccggtg ttaaaacttt 960 cgaattaaga actaaatcta aggaacaatt agaatctcaa ttggttgaat tgaaacaaga 1020 attggccact ttaaaagttc aaaaattaca aagaccaagt ttaccaagaa ttcacactgt 1080 tcgtaaaaac attgctagag tattgactgt tattaacttg aatcaaagag aaaatgttcg 1140 tgccttttac gctggtaaaa aatacattcc aaaagattta agagctaaaa agactagagc 1200 tttaagaaga aaattgacta aatttgaagc ttctcaagaa actgaaaaag ctagaaaaca 1260 1303 aagaattgct tttccacaaa gaaaatttgc tattaaagct taa

<210> 320

<211> 120

<212> PRT

<213> Candida albicans

<400> 320

Met Ala Gly Val Lys Thr Phe Glu Leu Arg Thr Lys Ser Lys Glu Gln
1 5 10 15

Leu Glu Ser Gln Leu Val Glu Leu Lys Gln Glu Leu Ala Thr Leu Lys
20 25 30

SEQL

```
Val Gln Lys Leu Gln Arg Pro Ser Leu Pro Arg Ile His Thr Val Arg
35 40 45
```

Lys Asn Ile Ala Arg Val Leu Thr Val Ile Asn Leu Asn Gln Arg Glu 50 55 60

Asn Val Arg Ala Phe Tyr Ala Gly Lys Lys Tyr Ile Pro Lys Asp Leu 65 70 75 80

Arg Ala Lys Lys Thr Arg Ala Leu Arg Arg Lys Leu Thr Lys Phe Glu 85 90 95

Ala Ser Gln Glu Thr Glu Lys Ala Arg Lys Gln Arg Ile Ala Phe Pro 100 105 110

Gln Arg Lys Phe Ala Ile Lys Ala 115 120

<210> 321

<211> 2690

<212> DNA

<213> Candida albicans

<400> 321

ctctgtgtaa attgatgaaa tccacacaat aaaaattttc tttcttcttt taagaaccta 60 aaaacagaat caacattatt tgccccatac atatccaaga attaaatact tattagttct 120 aagtggaata gaagagaatc aaacttaaca ttactgttac gcaacgtcaa gagggcattt 180 tttttagttt taatttgttt catttcaatt gaatctttaa gaatcaccga gtatacatac 240 tttcttttgt attttatcag ggaagccaca tccaaccacc agttacatcc cacaaaatcc 300 cttaatcttg ttcttagttg tattattaat ctattgaatt taagtttgat atgcgagaga 360 atattgtgga ttgtataagt tttgaactgg acttgaatac tttgaggggc ttaatcatat 420 attgcatttt ataccctact cgcggtgttg cttaccacac tgactagtat gatctttctg 480 agatttctag ccaataaatt atgagtgata tttatattat tattcatatt tctactacct 540 gtgatgactc gccgacattt gtcactaagg attcttccga gttaattgaa tttgcttggg 600 aaacagtcga tagtgtcact ttggaaactt tatacaaagg atcaaacttg gttcgtccaa 660 ccaacacacc tatcacacct tactgctcga agattcatag aataacatgg gacaatgtca 720 aaaatgctgg gtcgttcaaa gacgccatca caaactttga tcaatacgta caagaacaca 780 taatttccaa gaaaaaggag ttttcaattg tgatgtttga catttccaaa ttgagagttc 840 agttggttcg tgaagctaga gacaaatccg tggttttacc ctcgtatcta caacatccaa 900 ggatttttga tttaccaaga gaatatttaa attggcaatc tagccaccct gaaacattat 960 cataccccc aacttcttta actaatatta ttactgcatt agaagttgag gttgagaata 1020 tatctgaata tgtcgacttg ccaaactttt cttccacacc atcaccatca aaagcttcag 1080 caacaacgac gacgacaact gcaaatgtca cagccattga cgtcctttcc agtgaaacag 1140 aaccaaatgg taaagtcatt gcaaatttgc acgccaaaat tgccaaacaa ttgatcaaaa 1200 aatccatccc tgttgagaat caccctaatg tatttacaag accttttgat tcggctcaag 1260 atatcactgc ttttacatca gaaagatcaa aagtactcta tctttccaac ttgccaaacg 1320

```
acaccacaca atcagagetg gaatcatggt tcactcagta tggtggaaga ccaggtgggt 1380
tttggacttt taagtctgca gatgataaca ataataataa caacaacaat agcaatggcg 1440
ggaaaggata tcagaatgcg agaaaatatg gtatttcagg gtttgtggcc tttaatactc 1500
atgaggaagc agttgattgt ttagctttga atgggagagt gttgaatgat cgtcctattg 1560
aagttcaagc gtcttctagt aaagtgtttg atatggccat ggataaattg ttgttgactc 1620
tgttcccact ctcaaagaac agacctagac ccggggattg gacttgttta tcttgtgggt 1680
tttccaattt ccagagaaga acacactgtt tcaggtgctc ttttgcggca gtggcgtttc 1740
aggatgtttt taacagtaat acaggcaatg ccaacggtaa tggcaatgtt agcggcaacc 1800
acaaccacaa ccacaatagt ggagctcgcc gtggcatgaa tttacagcct gctcaagcta 1860
atgagaaaat tggaacaggc aatattagta ttccttctta caacgatcca atcaagggtc 1920
caacaggtaa tgtcactaat cacctcaaca attctgagac caatttactg aacaacacta 1980
atcttaacaa caacaatcat catagtaata attatcacaa taactaccat caccacaaca 2040
acaataataa caatcatggg aatagcaatg gtaacaccat acatggtcgt tcccattata 2100
acaatagtgt tccatttaga gcaggtgact ggaaatgtga aaattgcatg tatcacaatt 2160
tcgccaaaaa tttgtgttgt ttaaaatgtg gtgtcgccaa acctgctatt aacaatcaac 2220
aaaataatac aattcattcg gtgaattcaa cggccgctgc catagctgca gcaacagcca 2280
gtggtcaacc tttaaacttg aataataatg catttttgaa ccttcagcaa caacagtctc 2340
agtcacaacc ccaaggtcag caccattaca accaacattc tcgtaacaac aatgcttctg 2400
gggcatcaaa gttcaacaat ggctacaacc caaagaatca gtattacaat aataatagca 2460
agaatcttag caacaatttt ggtcttaatg gtatgcatca gcaaaaccaa aatcaaattt 2520
tgatgtattc acaacaattg caacaacaac agcaacaaca acagcaacaa cagcaacaac 2580
agcaacaaca gcaacaacag caacaacagc aacaacagca acagcaacaa catgatttaa 2640
atggaagtag ctcttcccat caactgaaac ttcaattgaa taatacttga
```

<210> 322

<211> 729

<212> PRT

<213> Candida albicans

<400> 322

Met Ser Asp Ile Tyr Ile Ile Ile His Ile Ser Thr Thr Cys Asp Asp

1 5 10 15

Ser Pro Thr Phe Val Thr Lys Asp Ser Ser Glu Leu Ile Glu Phe Ala 20 25 30

Trp Glu Thr Val Asp Ser Val Thr Leu Glu Thr Leu Tyr Lys Gly Ser
35 40 45

Asn Leu Val Arg Pro Thr Asn Thr Pro Ile Thr Pro Tyr Cys Ser Lys 50 55 60

Ile His Arg Ile Thr Trp Asp Asn Val Lys Asn Ala Gly Ser Phe Lys
65 70 75 80

Asp Ala Ile Thr Asn Phe Asp Gln Tyr Val Gln Glu His Ile Ile Ser 85 90 95

Lys	Lys	Lys	Glu 100	Phe	Ser	Ile	Val	Met 105	Pne	Asp	11e	ser	110	rea	Arg
Val	Gln	Leu 115	Val	Arg	Glu	Ala	Arg 120	Asp	Lys	Ser	Val	Val 125	Leu	Pro	Ser
Tyr	Leu 130	Gln	His	Pro	Arg	11e 135	Phe	Asp	Leu	Pro	Arg 140	Glu	Tyr	Leu	Asn
	Gln		Ser									Pro	Thr		Leu 160
Thr	Asn	Ile	Ile	Thr 165	Ala	Leu	Glu	Val	Glu 170	Val	Glu	Asn	Ile	Ser 175	Glu
Tyr	Val	Asp	Leu 180	Pro	Asn	Phe	Ser	Ser 185	Thr	Pro	Ser	Pro	Ser 190	Lys	Ala
Ser	Ala	Thr 195	Thr	Thr	Thr	Thr	Thr 200	Ala	Asn	Val	Thr	Ala 205	Ile	Asp	Val
Leu	Ser 210	Ser	Glu	Thr	Glu	Pro 215	Asn	Gly	Lys	Val	Ile 220	Ala	Asn	Leu ·	His
Ala 225	Lys	Ile	Ala	Lys	Gln 230	Leu	Ile	Lys	Lys	Ser 235	Ile	Pro	Val	Glu	Asn 240
His	Pro	Asn	Val	Phe 245	Thr	Arg	Pro	Phe	Asp 250	Ser	Ala	Gln	Asp	Ile 255	Thr
Ala	Phe	Thr	Ser 260	Glu	Arg	Ser	Lys	Val 265	Leu	Tyr	Leu	Ser	Asn 270	Leu	Pro
Asn	Asp	Thr 275	Thr	Gln	Ser	Glu	Leu 280		Ser			Thr 285	Gln	Tyr	Gly
Gly	Arg 290	Pro	Gly	Gly	Phe	Trp 295	Thr	Phe	Lys	Ser	Ala 300		Asp	Asn	Asn
Asn 305	Asn	Asn	Asn	Asn	Asn 310	Ser	Asn	Gly	Gly	Lys 315	Gly	Tyr	Gln	Asn	Ala 320
Arg	Lys	Tyr	Gly	11e 325	Ser	Gly	Phe	Val	Ala 330	Phe	Asn	Thr	His	G1u 335	
Ala	Val	Asp	Cys 340	Leu	Ala	Leu	Asn	Gly 345	Arg	Val	Leu	Asn	Asp 350	Arg	Pro

384



Ile Glu Val Gln Ala Ser Ser Ser Lys Val Phe Asp Met Ala Met Asp 355 360 365

Lys Leu Leu Leu Thr Ser Phe Pro Leu Ser Lys Asn Arg Pro Arg Pro 370 375 380

Gly Asp Trp Thr Cys Leu Ser Cys Gly Phe Ser Asn Phe Gln Arg Arg 385 390 395 400

Thr His Cys Phe Arg Cys Ser Phe Ala Ala Val Ala Phe Gln Asp Val 405 410 415

Phe Asn Ser Asn Thr Gly Asn Ala Asn Gly Asn Gly Asn Val Ser Gly 420 425 430

Asn His Asn His Asn His Asn Ser Gly Ala Arg Arg Gly Met Asn Leu 435 440 445

Gln Pro Ala Gln Ala Asn Glu Lys Ile Gly Thr Gly Asn Ile Ser Ile 450 455 460

Pro Ser Tyr Asn Asp Pro Ile Lys Gly Pro Thr Gly Asn Val Thr Asn 465 470 475 480

His Leu Asn Asn Ser Glu Thr Asn Leu Ser Asn Asn Thr Asn Leu Asn 485 490 495

Asn Asn Asn His His Ser Asn Asn Tyr His Asn Asn Tyr His His His 500 505 510

Asn Asn Asn Asn Asn His Gly Asn Ser Asn Gly Asn Thr Ile His 515 520 525

Gly Arg Ser His Tyr Asn Asn Ser Val Pro Phe Arg Ala Gly Asp Trp 530 535 540

Lys Cys Glu Asn Cys Met Tyr His Asn Phe Ala Lys Asn Leu Cys Cys 545 550 555 560

Leu Lys Cys Gly Val Ala Lys Pro Ala Ile Asn Asn Gln Gln Asn Asn 565 570 575

Thr Ile His Ser Val Asn Ser Thr Ala Ala Ala Ile Ala Ala Ala Thr 580 585 590

Ala Ser Gly Gln Pro Leu Asn Leu Asn Asn Asn Ala Phe Leu Asn Leu 595 600 605

```
Gln Gln Gln Gln Ser Gln Ser Gln Pro Gln Gly Gln His His Tyr Asn 610 615 620 620

Gln His Ser Arg Asn Asn Asn Ala Ser Gly Ala Ser Lys Phe Asn Asn 625 630 635 640

Gly Tyr Asn Pro Lys Asn Gln Tyr Tyr Asn Asn Asn Ser Lys Asn Leu 645 650 655

Ser Asn Asn Phe Gly Leu Asn Gly Met His Gln Gln Asn Gln Asn Gln Asn Gln 660 665
```

Gln Gln Gln Gln Gln His Asp Leu Asn Gly Ser Ser Ser His
705 710 715 720

Gln Ser Lys Leu Gln Leu Asn Asn Thr 725

<210> 323 <211> 3359 <212> DNA <213> Candida albicans

## <400> 323

aatagcattg aacaagaaga agaggacaac gatagaccaa ggttggtttt agccaatcct 60 gattatgata gtgatgacag ttcatagaca atttacagcc ttaaatggat atatatgtat 120 atttaataat aaaggacttg tttttttag taactagtgt gatctctttt ctgggtgtac 180 atttcggata gccaaccagg ttatatttta gcagtttata gacagtgtta tcgatgggta 240 atataaataa aagctcattg aatactatct agtgaaaagt cgtgtgtaaa tcgatttgaa 300 aacaaaaaaa ccatacacgt aaatgaaatg tgtgtgaaag tacaaccaac aacgaaaaag 360 aacaaaaaaa aaatgttgtc cgccaaaaaa aaaaagagaa caacaaatca aagtttcaag 420 actatctcaa atcttgttgt caccataact atcaattgtt cacctcttga accaacatca 480 cggctacgaa agtatctgtc aatgagtgaca gtggttatac attaatctat gagcctaata 540 cggctacgaa agtatctgtc aatgaatta aaaatttgtt ggaaaagggt aaagatgatg 600 tgaaagtaga taccatgaag aagattttga ttaccatatt aaatggagac cccttacctg 660 acttgttgat gcatataatc agatttgca tgcctccag aaataaagaa ttgaaaaagt 720 tgttgtatca ttattgggag gtttgtccaa aaatggatga atcaggtaaa atgagacatg 780 aaatgattct tgtgtgtaat gccatccaac gtgatttaca gcatccaaat gaatatattc 840 gaggcaatac tttgaggatat ttgacgaaat tgaaagagcc agaattattg gaaactttag 900

```
ttcctaatgt ccgtcaatgt ttagaacacc gtcatgccta tgtcagaaaa aatgctgttt 960
tcgcattatg gtctattcat aaagtcagtg atcatttagc tcctgatgct gacgagttaa 1020
tttacagatt tttgtatgag gaaaacgatt ctgtttgtaa aagaaatgct tttgtttgtc 1080
ttggagactt gaatagagaa gctgctttgc aatatattca ggataatatt tcagttattg 1140
agactttgga tccattgata caattggctt ttattgagtt tatcaaaaag gactctattc 1200
aaaatccagc tttaaagcaa caatatgccc aattaatgac agaaattatt gaaagctctt 1260
caaatgttgt tatgtatgaa gctgctaaca cgttgactgt tttgacttca aacccacaat 1320
caattttgtt ggcaggaaac aagtttgttg aattggctac tagagagtct gataataacg 1380
ttaaaattat cactttagag agaataaatc aattacacaa gcaacatcct ggtgtgttac 1440
aagacttgtc attagaaatt ttacgaggtt tatcttccca agatttggat gttaaaaaga 1500
aagctcttga tgttacttta caatttatca ccaccagaaa tgttgaagat gttgttaagt 1560
tattgaagaa agaattgcag tctacagctt tatccaatga tgacaagaat gcagattata 1620
gacagttgtt aattaatgcc atccatcaat tggctattaa atttgtggag gttgctgcca 1680
atgtcattga tttattgttg gattctatag ccgatttgaa taccactgcc gcctacgagg 1740
ttatcacatt tgttaaagaa gttgttgaga aattcccaga tttaagggac gctattttga 1800
gaagattgat tttggctttg ccacatgtga aaagtggtaa agttttccgt ggtgcattat 1860
gggttattgg tgagtatgcc ttagaggaat cattaataca agaatcttgg aaatatatta 1920
gaggaagtat tggtgaagta ccaattattg ctagtgaatt gaaactgaaa aagcgtgatg 1980
atactgagga atcacaagag gaagaaaccg agtatgatgg taaacctcgc agaaagggtc 2040
cagttqtatt gccagatggt acctacgcta ctgagtcagc attgacaagc gaaacaactg 2100
actctttgga aagtgacagc aagactccta tcagaaagca aattcttgct ggtgatttct 2160
acttaggtgc tgtattagca tcaactttgg tgaaattgat tctccgattg caaagtttga 2220
aacaaactca agaaaaaatt ttgaacggat taaaagcaga agcattgttg attatggttt 2280
cgattttaag agttggggaa tctagcttgg tttctaagaa aattgatgag gattctgctg 2340
acagaatttt gtcttacatc aagattttga acgatgaaga agaccttcag gaaatcaaga 2400
caagetteet tgaagataet aaagatgeat ttaaageaca aattaataat getgaattga 2460
agaaagcaga agcattggct aaggatttgc atgataacgc tgaacaaatt gacgatgcaa 2520
ttgtttttag acagttggat aaagataaca aaaagagtaa agcttctgtg gatgatgttg 2580
ctgctgcgtc aggaagcaat gaattaaaga aagaaaattt gtcgtcgaga ttgaacaaaa 2640
ttatacaatt gactgggttt tccgatccta tttacgcaga ggcatttgtc aaagttcatc 2700
aatacgatgt tgtgttagat gtcttgctag tgaatcaaac cacaactact ttaagaaact 2760
tatcagttga atttgctaca ttgggtgatt tgaaagtggt tgataaacca actaccgcaa 2820
atattggacc tcatggtttc tacaaagttc aaacaactat taaagttact tcggctgata 2880
ctggtgtcat ctttggtaac atagtgtatg acggtcaaca ctcggacgat tcacgtatag 2940
ttattttgaa tgacgttcac gttgacatta tggattacat taagccagcc acttgttcag 3000
aaagtcaatt ccgtaaaatg tggaacgaat ttgaatggga gaataagata accattaaat 3060
cacctattga aacattgaaa gagtacttgg atgaattaat gaagggtaca aatatgcaat 3120
gcttgacacc gggtgcggta attggagaag aatgtcaatt tttatcagca aacttgtact 3180
caaggtcaag ctttggtgaa gatgcattgg ctaatttatg tatagagaaa cagagtgatg 3240
gaccaataat tggtcatgtc agaataagat caaaaggtca aggtttggct ttgtcattgg 3300
gtgatagagt agcttccatt tcaagaaaag gtaagaaggc aactattgct cgtgtttaa 3359
```

<sup>&</sup>lt;210> 324

<sup>&</sup>lt;211> 952

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Candida albicans

<400>	32	4
-------	----	---

Met Ser Asp Ser Gly Tyr Thr Leu Ile Tyr Glu Pro Asn Thr Ala Thr 1 5 10 15

Lys Val Ser Val Asn Glu Phe Lys Asn Leu Leu Glu Lys Gly Lys Asp 20 25 30

Asp Val Lys Val Asp Thr Met Lys Lys Ile Leu Ile Thr Ile Leu Asn 35 40 45

Gly Asp Pro Leu Pro Asp Leu Leu Met His Ile Ile Arg Phe Val Met

50 55 60 60

50 55 60

Pro Ser Arg Asn Lys Glu Leu Lys Lys Leu Leu Tyr His Tyr Trp Glu 65 70 75 80

Val Cys Pro Lys Met Asp Glu Ser Gly Lys Met Arg His Glu Met Ile 85 90 95

Leu Val Cys Asn Ala Ile Gln Arg Asp Leu Gln His Pro Asn Glu Tyr
100 105 110

Ile Arg Gly Asn Thr Leu Arg Tyr Leu Thr Lys Leu Lys Glu Pro Glu
115 120 125

Leu Leu Glu Thr Leu Val Pro Asn Val Arg Gln Cys Leu Glu His Arg 130 135 140

His Ala Tyr Val Arg Lys Asn Ala Val Phe Ala Leu Trp Ser Ile His 145 150 155 160

Lys Val Ser Asp His Leu Ala Pro Asp Ala Asp Glu Leu Ile Tyr Arg 165 170 175

Phe Leu Tyr Glu Glu Asn Asp Ser Val Cys Lys Arg Asn Ala Phe Val 180 185 190

Cys Leu Gly Asp Leu Asn Arg Glu Ala Ala Leu Gln Tyr Ile Gln Asp 195 200 205

Asn Ile Ser Val Ile Glu Thr Leu Asp Pro Leu Ile Gln Leu Ala Phe 210 215 220

Ile Glu Phe Ile Lys Lys Asp Ser Ile Gln Asn Pro Ala Leu Lys Gln 225 230 235 240

Gln Tyr Ala Gln Leu Met Thr Glu Ile Ile Glu Ser Ser Ser Asn Val 245 250 255

Val Met Tyr Glu Ala Ala Asn Thr Leu Thr Val Leu Thr Ser Asn Pro 260 265 270

Gln Ser Ile Leu Leu Ala Gly Asn Lys Phe Val Glu Leu Ala Thr Arg 275 280 285

Glu Ser Asp Asn Asn Val Lys Ile Ile Thr Leu Glu Arg Ile Asn Gln 290 295 300

Leu His Lys Gln His Pro Gly Val Leu Gln Asp Leu Ser Leu Glu Ile 305 310 315 320

Leu Arg Gly Leu Ser Ser Gln Asp Leu Asp Val Lys Lys Lys Ala Leu 325 330 335

Asp Val Thr Leu Gln Phe Ile Thr Thr Arg Asn Val Glu Asp Val Val 340 345 350

Lys Leu Leu Lys Lys Glu Leu Gln Ser Thr Ala Leu Ser Asn Asp Asp 355 360 365

Lys Asn Ala Asp Tyr Arg Gln Leu Leu Ile Asn Ala Ile His Gln Leu 370 375 380

Ala Ile Lys Phe Val Glu Val Ala Ala Asn Val Ile Asp Leu Leu Leu 385 390 395 400

Asp Ser Ile Ala Asp Leu Asn Thr Thr Ala Ala Tyr Glu Val Ile Thr 405 410 415

Phe Val Lys Glu Val Val Glu Lys Phe Pro Asp Leu Arg Asp Ala Ile 420 425 430

Leu Arg Arg Leu Ile Leu Ala Leu Pro His Val Lys Ser Gly Lys Val
435
440
445

Phe Arg Gly Ala Leu Trp Val Ile Gly Glu Tyr Ala Leu Glu Glu Ser 450 455 460

Leu Ile Gln Glu Ser Trp Lys Tyr Ile Arg Gly Ser Ile Gly Glu Val 465 470 475 480

Pro Ile Ile Ala Ser Glu Leu Lys Ser Lys Lys Arg Asp Asp Thr Glu 485 490 495

Glu Ser Gln Glu Glu Glu Thr Glu Tyr Asp Gly Lys Pro Arg Arg Lys
500 505 510

Gly Pro Val Val Leu Pro Asp Gly Thr Tyr Ala Thr Glu Ser Ala Leu Thr Ser Glu Thr Thr Asp Ser Leu Glu Ser Asp Ser Lys Thr Pro Ile Arg Lys Gln Ile Leu Ala Gly Asp Phe Tyr Leu Gly Ala Val Leu Ala Ser Thr Leu Val Lys Leu Ile Leu Arg Leu Gln Ser Leu Lys Gln Thr <del>--565--</del> Gln Glu Lys Ile Leu Asn Gly Leu Lys Ala Glu Ala Leu Leu Ile Met Val Ser Ile Leu Arg Val Gly Glu Ser Ser Leu Val Ser Lys Lys Ile Asp Glu Asp Ser Ala Asp Arg Ile Leu Ser Tyr Ile Lys Ile Leu Asn Asp Glu Glu Asp Leu Gln Glu Ile Lys Thr Ser Phe Leu Glu Asp Thr 

Lys Asp Ala Phe Lys Ala Gln Ile Asn Asn Ala Glu Leu Lys Lys Ala 

Glu Ala Leu Ala Lys Asp Leu His Asp Asn Ala Glu Gln Ile Asp Asp 

Ala Ile Val Phe Arg Gln Leu Asp Lys Asp Asn Lys Lys Ser Lys Ala 

Ser Val Asp Asp Val Ala Ala Ala Ser Gly Ser Asn Glu Leu Lys Lys 

Glu Asn Leu Ser Ser Arg Leu Asn Lys Ile Ile Gln Leu Thr Gly Phe 

Ser Asp Pro Ile Tyr Ala Glu Ala Phe Val Lys Val His Gln Tyr Asp 

Val Val Leu Asp Val Leu Leu Val Asn Gln Thr Thr Thr Thr Leu Arg 

Asn Leu Ser Val Glu Phe Ala Thr Leu Gly Asp Leu Lys Val Val Asp 

Lys Pro Thr Thr Ala Asn Ile Gly Pro His Gly Phe Tyr Lys Val Gln 770 775 780

Thr Thr Ile Lys Val Thr Ser Ala Asp Thr Gly Val Ile Phe Gly Asn 785 790 795 800

Ile Val Tyr Asp Gly Gln His Ser Asp Asp Ser Arg Ile Val Ile Leu 805 810 815

Asn Asp Val His Val Asp Ile Met Asp Tyr Ile Lys Pro Ala Thr Cys
820 825 830

Ser Glu Ser Gln Phe Arg Lys Met Trp Asn Glu Phe Glu Trp Glu Asn 835 840 845

Lys Ile Thr Ile Lys Ser Pro Ile Glu Thr Leu Lys Glu Tyr Leu Asp 850 855 860

Glu Leu Met Lys Gly Thr Asn Met Gln Cys Leu Thr Pro Gly Ala Val 865 870 875 880

Ile Gly Glu Cys Gln Phe Leu Ser Ala Asn Leu Tyr Ser Arg Ser 885 890 895

Ser Phe Gly Glu Asp Ala Leu Ala Asn Leu Cys Ile Glu Lys Gln Ser 900 905 910

Asp Gly Pro Ile Ile Gly His Val Arg Ile Arg Ser Lys Gly Gln Gly 915 920 925

Leu Ala Leu Ser Leu Gly Asp Arg Val Ala Ser Ile Ser Arg Lys Gly 930 935 940

Lys Lys Ala Thr Ile Ala Arg Val 945 950

<210> 325

<211> 2270

<212> DNA

<213> Candida albicans

<400> 325

gtttgatcac atgtgttgtg aacactcggg taatacaaaa tagtgagaga gaagaagggg 60 aaaaaaaaaa gcagaacaca aaacatggaa tttgaaaaca attttgtaat tcaatcgatc 120 cgagacttcc atagcaaagt taacaagcac aattgtcatt ttaacttaat tggtggatta 180

```
atcgggatca atctgaattg tttcccgtat tgtttaaaac caaagaaaaa ggataatcaa 240
aactaaatct ttcatattaa cactaccatt ttgtagtggt cagtttatat aattatccac 300
tgttctcttc acaattaaaa aaaaaaaaaa aaagaaaaaa gaaagctctc tctcccccaa 360
aagaaaaagc aaaggtaatt ccttcataca cacctttgat atctttcctc ttagactttt 420
cttttaatac ttgcatcaat tggaatatta cttgttcata ctggagtttt cattgaacta 480
aatattatta aaatattatt atgettgaat tgaatteaat taegatteet eggaatttea 540
ctgaattcca actcacagca ttaaagatct actatcaact taagatttta tttcttgcca 600
catactgtgc tcaaggatca tttggcttga acggttcagt ctgtttagca agagatattt 660
ttgttggcta tgttgtatac actcaattat taaagcttta tcgagtatta agagggtacg 720
gtattgtaga ttctattcga aggttatacc tatatgttag ttctacggtg tcatctcaaa 780
tetttteact accatttatt aaateeaaaa ttgacaagga attgeaageg actattggea 840
aagtagaaga agagattatg aaaaacgatc cacagttatt acagtttccc gaattgccag 900
aacagggtat tgacgctgac aatgtttcct tggagttgga taaattacaa aacttgaaac 960
attctgactg gatcaatgga agagtcagtg gagcagttta tcatggaggt gagaatctat 1020
tgtcattaca agttgaagct tacaagaaat actcggtggc caatcaattg catcccgatg 1080
ttttcccagg tgtgcgtaag atggaggccg aagtagttca tatggttttg gatatcttta 1140
atgctccgag tgacgggtgt ggatcgacaa cttcgggagg tactgagtcg ttattgttag 1200
ccgggttgtc agctagagaa tatgggaaga aatatcgtgg aataactgag cctgaagtca 1260
ttgcccccgt gacaatacat gcgggaattg aaaaagcctg tttttatttt ggaatgaaat 1320
tacataaagt tgacttagat ccggttacct ttcaagttga cgttaagaaa gtagaaagat 1380
tgatcaatag taatacagtt ttaatctgtg gatcagcccc aaattatcct catggaataa 1440
ttgatgatat agagtcctta tccaagttgg cagtcaagta taatatcccg ttgcacgttg 1500
atgcatgttt ggggtcattt attgtttcgt ttttagaaaa atcaaaagta catggcgata 1560
ggaaattgcc catatttgat tttcgattac caggtgtcac gtcaatctca tgtgatactc 1620
acaaatatgg gtttgctccc aaggggtcat caataattat gtaccgttcg ccaaaattac 1680
gtgagtgtca atactatatt gcaagtgatt ggacgggtgg aatgtatggt tctccaactt 1740
tggctggttc taggccaggt gctcttgtag ttggatgctg ggctacatta atcaatattg 1800
ggaaacaagg atacaccaag ttttgttacg atattgtgct ggcgtcaatg aaagttaaac 1860
gagcaattga aactgacccg atactatcca aacatttaca aattattggt gatccaattg 1920
ggtcggtaat ttcgtttcaa cttgcacctc agcaatcggg aaatttaagt atttacgaga 1980
ttagtgattt gttgaccaaa aaaggttggc attttgcaac tttacaaaac ccatcagcat 2040
tacattttgc atttacaaga ttgactgtcc cggtggtcga tgaattgatt gcagatttgg 2100
ttgaagctac aaaagaagcg gtggcaattg ccgaggaaca caaaaagaat ggagtgacca 2160
aagctccagg tgatactgct gcgttatatg gtatagctgg cagtgtacat acagcagggt 2220
tggctgatag attaattgtt gcatttttag ataccttata caaaatttga
                                                                  2270
```

```
<210> 326
```

## <400> 326

```
Met Leu Glu Leu Asn Ser Ile Thr Ile Pro Arg Asn Phe Thr Glu Phe 1 5 10 15
```

```
Gln Leu Thr Ala Leu Lys Ile Tyr Tyr Gln Leu Lys Ile Leu Phe Leu 20 25 30
```

<sup>&</sup>lt;211> 589

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Candida albicans

Ala Thr Tyr Cys Ala Gln Gly Ser Phe Gly Leu Asn Gly Ser Val Cys 35 40 45

1

Leu Ala Arg Asp Ile Phe Val Gly Tyr Val Val Tyr Thr Gln Leu Leu 50 55 60

Lys Leu Tyr Arg Val Leu Arg Gly Tyr Gly Ile Val Asp Ser Ile Arg 65 70 75 80

Arg Leu Tyr Leu Tyr Val Ser Ser Thr Val Ser Ser Gln Ile Phe Ser 85 90 95

Leu Pro Phe Ile Lys Ser Lys Ile Asp Lys Glu Leu Gln Ala Thr Ile 100 105 110

Gly Lys Val Glu Glu Glu Ile Met Lys Asn Asp Pro Gln Leu Leu Gln 115 120 125

Phe Pro Glu Leu Pro Glu Gln Gly Ile Asp Ala Asp Asn Val Ser Leu 130 135 140

Glu Leu Asp Lys Leu Gln Asn Leu Lys His Ser Asp Trp Ile Asn Gly
145 150 155 160

Arg Val Ser Gly Ala Val Tyr His Gly Gly Glu Asn Leu Leu Ser Leu 165 170 175

Gln Val Glu Ala Tyr Lys Lys Tyr Ser Val Ala Asn Gln Leu His Pro 180 185 190

Asp Val Phe Pro Gly Val Arg Lys Met Glu Ala Glu Val Val His Met 195 200 205

Val Leu Asp Ile Phe Asn Ala Pro Ser Asp Gly Cys Gly Ser Thr Thr 210 215 220

Ser Gly Gly Thr Glu Ser Leu Leu Leu Ala Gly Leu Ser Ala Arg Glu 225 230 235 240

Tyr Gly Lys Lys Tyr Arg Gly Ile Thr Glu Pro Glu Val Ile Ala Pro 245 250 255

Val Thr Ile His Ala Gly Ile Glu Lys Ala Cys Phe Tyr Phe Gly Met 260 265 270

Lys Leu His Lys Val Asp Leu Asp Pro Val Thr Phe Gln Val Asp Val 275 280 285

Lys	Lys 290	Val	Glu	Arg	Leu	Ile 295	Asn	Ser	Asn	Thr	Val 300	Leu	Ile	Cys	Gly
Ser 305	Ala	Pro	Asn	Tyr	Pro 310	His	Gly	Ile	Ile	Asp 315	Asp	Ile	Glu	Ser	Leu 320
Ser	Lys	Leu	Ala	Val 325	Lys	Tyr	Asn	Ile	Pro 330	Leu	His	Val	Asp	Ala 335	Суѕ
Leu	Gly	Ser	Phe	Ile	Val	Ser	Phe	Leu 345	Glu	Lys	Ser	Lys	Val 350	His	Gly
										,*					
Asp	Arg	Lys 355	Leu	Pro	Ile	Phe	360	Phe	Arg	Leu	Pro	Gly 365	Val	Thr	Ser
Ile	Ser 370	Cys	Asp	Thr	His	Lys 375	Tyr	Gly	Phe	Ala	Pro 380	Lys	Gly	Ser	Ser
Ile	Ile	Met	Tyr	Arg	Ser	Pro	Lys	Leu	Arg	Glu	Cys	Gln	Tyr	Tyr	Ile
385			-		390					395					400
Ala	Ser	Asp	Trp	Thr 405	Gly	Gly	Met	Туr	Gly 410	Ser	Pro	Thr	Leu	Ala 415	Gly
Ser	Arg	Pro	Gly 420	Ala	Leu	Val	Val	Gly 425	Cys	Trp	Ala	Thr	Leu 430	Ile	Asn
Ile	Gly	Lys 435	Gln	Gly	Tyr	Thr	Lys 440	Phe	Cys	Tyr	Asp	Ile 445	Val	Ser	Ala
Ser	Met 450	Lys	Val	Lys	Arg	Ala 455	Ile	Glu	Thr	Asp	Pro 460	Ile	Leu	Ser	Lys
His 465	Leu	Gln	Ile	Ile	Gly 470	Asp	Pro	Ile	Gly	Ser 475	Val	Ile	Ser	Phe	Gln 480
Leu	Ala	Pro	Gln	Gln 485	Ser	Gly	Asn	Leu	Ser 490	Ile	Tyr	Glu	Ile	Ser 495	Asp
Leu	Leu	Thr	Lys 500	Lys	Gly	Trp	His	Phe 505	Ala	Thr	Leu	Gln	Asn 510	Pro	Ser
Ala	Leu	His 515	Phe	Ala	Phe	Thr	Arg 520	Leu	Thr	Val	Pro	Val 525	Val	Asp	Glu
Leu	Ile 530	Ala	Asp	Leu	Val	Glu 535	Ala	Thr	Lys	Glu	Ala 540	Val	Ala	Ile	Ala

```
Glu Glu His Lys Lys Asn Gly Val Thr Lys Ala Pro Gly Asp Thr Ala
545 550 555 560
```

Ala Leu Tyr Gly Ile Ala Gly Ser Val His Thr Ala Gly Leu Ala Asp 565 570 575

Arg Leu Ile Val Ala Phe Leu Asp Thr Leu Tyr Lys Ile 580 585

```
<210> 327
<211> 3605
<212> DNA
<213> Candida albicans
```

## <400> 327

```
ttcaattttt ttaaaaaaat agcaaaaagg tatttcttag aaattgaaaa aaaaaaatca 60
caaaaaaatt tataaaaccg gatagggccg ttatcgcagg acgtgtccca tgatcaatct 120
acaatgaaat gattcacatg aaatatgtta caattccaca cttgctgcta aaaagcaaca 180
gttttgcaat agagaaaagc atgatatact atcgataata tctttcgata agaacttaaa 240
tgtagcaacg ttgaatttta aaataaaact tccttttcct ggtgataaat tttgactttg 300
aagcataaag aagacagagc taaaaaaaaa ctactctcta gacaaaatca aaacctcgaa 360
tatatttttg gtgtttcttt ttttttttt ttgcctctcg ttgaaatcat ctccattctt 420
cttttccacg tctttgttga aaactttgca acctaaaaaa ataagaatca ctccaacaag 480
tttaaactac attatcaatt atgttgaaaa ctagattaaa acaaagcagg gccataagtc 540
gggttgtaag aagatatgca tgctcacacc ccatttctcc caatcttgat aagtacccag 600
ttggtctaaa attgcatggt tacgaagtta cccaaacatc acctatccca gaattttccc 660
tcactgctgt atcattaaaa cacacagaga gtggtgcaac tcacttacat ttggattccc 720
ctaatgacag taataatgta tttctgattg ccttcaaaac aaatcctcca gataatactg 780
gggttcccca tattttagaa catacaactt tgtgtggtag taaaaagttt ccggtccgtg 840
atccattttt taaaatgacc aacaggtcgt tgagtaactt tatgaatgca atgacaggcc 900
atgattacac attttatcca tttgctacca ccaattcaaa ggattttgaa aacctaatgg 960
atgtgtattt atcgtcagtg tttgaaccgc aattaaacca taccgatttc ttgcaagaag 1020
gatggagaat agaaaatcaa aatgttcatg acatatcgtc caagcttgaa ttcaagggag 1080
ttgtatataa tgaaatgaag ggccagtatt cgaactctgc atactacttt tatatcaagt 1140
tccttgagag tatatatcca tccttgaata attcaggggg tgatcccaag aaaattgttg 1200
atttgctgta cgagggttta ctagagtttc acctgaaaaa ttatcatcca tcaaatgcaa 1260
aaacatttac ttatggaaaa ttaccattgg aagacagttt aagtaaaata agcaaatact 1320
atgaatcatt cgaaaaaaag gtgtcttcag ttgacgtcaa acaacctata ttttctacag 1380
ataaatcaga aatctttgat gtcaccatcc cgggtccagt tgatacaatg aatggtaaag 1440
agacttcaga acagtactgc acatctatca cctggaactt gggtaatcca ttggatccaa 1500
acatgcagta tgatattttt aaatggaaaa tattgagctc attattgttt gatggacaca 1560
actctccttt ctatcaagag ttaattgaaa gtggatacgg tgacgatttt tctgcaaata 1620
ctgggttgga ctcaaccacc gcgttgcttt cgtttactgt tggtctcaac tacttaacca 1680
agcaaaaagt tgataatttt aatgaaaaag ttatggaaat cattaataat aaaatcattc 1740
ccgaattaag taacgaagag tcctcttcat atcatggtag aattgatgct atattgcatc 1800
```

```
aaatagaaat aggattcaaa agacacaagc ccgattttgg atttggatta ttgagctcta 1860
ttgttccgtc atgggtgaat ggagttgatc caattgacac cttgcaagtg gaaaagatat 1920
tgtcgcattt taaagaagat tataaacaaa atggtttaag gatctttaaa gaattattag 1980
aaaagacatt gtgtaaccct cattcgcaaa aatttaaatt caccatggag ccaagagaag 2040
attttaccaa acaattggta aaagatgaga atttgatgat cgagaaaaga gtaagtgaac 2100
tcacagaaga taacaagaag gcaatctatg agcaaaactt ggaattagct aaattacaat 2160
tggaggatca aaatacagaa gttttaccca cattgactat tgatgacatt ccaaagagag 2220
gtgattttta tgccattgat ttgggccaag taaataagaa agttgtacat gaaagggtag 2280
ttgataccaa tggcttggtt tatgccaacg ctttaaaaaga tatttcctat ttacccacca 2340
aactttacaa gtaccttcca ttgtttaaca actgtttgac gaaccttgct ggaacagaaa 2400
acacacccat tacggagttg gaaactaaaa tacaaatgtt aactggcggg ataacattta 2460
gttctaaaat atcgactgac ccctataata ttgagcaact aaaattacag tatgtgttaa 2520
gtggaatggc tttgaaagaa aagtcatcct cagtttatga tttatggttg gagattttaa 2580
ctactaccaa attcgacacc agtgatgagg tattagaaaa gttgtcagtt ttgattaaaa 2640
acatgggaca aaaccaaatc aataatattg ctgatcgcgg tcattcttat gcggctgctg 2700
tgagctcact gaaattgaca ccgctgaaat acatcagtga catcgtttca ggtttgagtc 2760
aagttcaatt tgtaatggag ttgaactcca aattagaatc agaagggaaa gagtacttgg 2820
ccaaagagat tattccgata ttgcaagaaa tacaaaagta tgtattgcaa ggtgaattca 2880
ggtatagact agttggaaat caagagatta ttgttgaaaa cgaaaagctt attgagaaat 2940
ttgataagga tatttcttcg aacagaccaa ctttatcgtt aacagtaaca gatggtttac 3000
tggcattgtt gaactcattc aattacaatc atacaagtga aaatgtctta gttaacttac 3060
catttcaagt gggatactct tcattaggta agattggctc ttcgtattca tcaaaggatg 3120
gtgcttcttt acaaatatta tctcagttat attcctttaa aaatctacat tccaaaataa 3180
gagaaagcaa tggtgcatat ggaggtggtt tgacatatga tgggttgaac gggacattaa 3240
actittattc gtatcgtgat cctaatcctg ttaagtcgat tcaaactttt agagattcct 3300
tactgtatgg acttgatgct aattggaacg ataaggattt acaagaggct aagttgcggg 3360
ttttccaaag cgtcgatgct ccaattaata tctcttctca gggtgctagt gccttctttg 3420
aaaatataga tgattacttg agacaggaaa gaagagaaaa ctttttgggt accactttaa 3480
aggateteag agatgtgaet gaaaagtate ttgttgataa ccaaaacaac ettgtcaetg 3540
ttattggtga caatgaaatt ttaaatgtcg ataataaatg gcaaattaga aattttcaag 3600
                                                                  3605
tatag
```

```
<210> 328
<211> 1034
<212> PRT
```

<213> Candida albicans

<400> 328

Met Leu Lys Thr Arg Leu Lys Gln Ser Arg Ala Ile Ser Arg Val Val
1 5 10 15

Arg Arg Tyr Ala Cys Ser His Pro Ile Ser Pro Asn Leu Asp Lys Tyr
20 25 30

Pro Val Gly Leu Lys Leu His Gly Tyr Glu Val Thr Gln Thr Ser Pro 35 40 45 Ile Pro Glu Phe Ser Leu Thr Ala Val Ser Leu Lys His Thr Glu Ser
50 55 60

Gly Ala Thr His Leu His Leu Asp Ser Pro Asn Asp Ser Asn Asn Val 65 70 75 80

Phe Ser Ile Ala Phe Lys Thr Asn Pro Pro Asp Asn Thr Gly Val Pro 85 90 95

His Ile Leu Glu His Thr Thr Leu Cys Gly Ser Lys Lys Phe Pro Val

Arg Asp Pro Phe Phe Lys Met Thr Asn Arg Ser Leu Ser Asn Phe Met 115 120 125

Asn Ala Met Thr Gly His Asp Tyr Thr Phe Tyr Pro Phe Ala Thr Thr 130 135 140

Asn Ser Lys Asp Phe Glu Asn Leu Met Asp Val Tyr Leu Ser Ser Val 145 150 155 160

Phe Glu Pro Gln Leu Asn His Thr Asp Phe Leu Gln Glu Gly Trp Arg 165 170 175

Ile Glu Asn Gln Asn Val His Asp Ile Ser Ser Lys Leu Glu Phe Lys 180 185 190

Gly Val Val Tyr Asn Glu Met Lys Gly Gln Tyr Ser Asn Ser Ala Tyr 195 200 205

Tyr Phe Tyr Ile Lys Phe Leu Glu Ser Ile Tyr Pro Ser Leu Asn Asn 210 215 220

Ser Gly Gly Asp Pro Lys Lys Ile Val Asp Leu Ser Tyr Glu Gly Leu 225 230 235 240

Leu Glu Phe His Ser Lys Asn Tyr His Pro Ser Asn Ala Lys Thr Phe 245 250 255

Thr Tyr Gly Lys Leu Pro Leu Glu Asp Ser Leu Ser Lys Ile Ser Lys 260 265 270

Tyr Tyr Glu Ser Phe Glu Lys Lys Val Ser Ser Val Asp Val Lys Gln 275 280 285

Pro Ile Phe Ser Thr Asp Lys Ser Glu Ile Phe Asp Val Thr Ile Pro 290 295 300

Gly Pro Val Asp Thr Met Asn Gly Lys Glu Thr Ser Glu Gln Tyr Cys Thr Ser Ile Thr Trp Asn Leu Gly Asn Pro Leu Asp Pro Asn Met Gln Tyr Asp Ile Phe Lys Trp Lys Ile Leu Ser Ser Leu Leu Phe Asp Gly His Asn Ser Pro Phe Tyr Gln Glu Leu Ile Glu Ser Gly Tyr Gly Asp Asp Phe Ser Ala Asn Thr Gly Leu Asp Ser Thr Thr Ala Leu Leu Ser Phe Thr Val Gly Leu Asn Tyr Leu Thr Lys Gln Lys Val Asp Asn Phe Asn Glu Lys Val Met Glu Ile Ile Asn Asn Lys Ile Ile Pro Glu Leu Ser Asn Glu Glu Ser Ser Tyr His Gly Arg Ile Asp Ala Ile Leu His Gln Ile Glu Ile Gly Phe Lys Arg His Lys Pro Asp Phe Gly Phe Gly Leu Leu Ser Ser Ile Val Pro Ser Trp Val Asn Gly Val Asp Pro Ile Asp Thr Leu Gln Val Glu Lys Ile Leu Ser His Phe Lys Glu Asp Tyr Lys Gln Asn Gly Leu Arg Ile Phe Lys Glu Leu Leu Glu Lys Thr Leu Cys Asn Pro His Ser Gln Lys Phe Lys Phe Thr Met Glu Pro Arg Glu Asp Phe Thr Lys Gln Leu Val Lys Asp Glu Asn Leu Met Ile Glu Lys Arg Val Ser Glu Leu Thr Glu Asp Asn Lys Lys Ala Ile Tyr Glu Gln Asn Leu Glu Leu Ala Lys Leu Gln Leu Glu Asp Gln Asn Thr Glu

Val Leu Pro Thr La Thr Ile Asp Asp Ile Pro Lys Arg Sly Asp Phe 565 570 575

Tyr Ala Ile Asp Leu Gly Gln Val Asn Lys Lys Val Val His Glu Arg 580 585 590

Val Val Asp Thr Asn Gly Leu Val Tyr Ala Asn Ala Leu Lys Asp Ile 595 600 605

Ser Tyr Leu Pro Thr Lys Leu Tyr Lys Tyr Leu Pro Leu Phe Asn Asn 610 615 620

Cys Leu Thr Asn Leu Ala Gly Thr Glu Asn Thr Pro Ile Thr Glu Leu 625 630 635 640

Glu Thr Lys Ile Gln Met Leu Thr Gly Gly Ile Thr Phe Ser Ser Lys 645 650 655

Ile Ser Thr Asp Pro Tyr Asn Ile Glu Gln Leu Lys Leu Gln Tyr Val 660 665 670

Leu Ser Gly Met Ala Leu Lys Glu Lys Ser Ser Ser Val Tyr Asp Leu 675 680 685

Trp Leu Glu Ile Leu Thr Thr Lys Phe Asp Thr Ser Asp Glu Val 690 695 700

Leu Glu Lys Leu Ser Val Leu Ile Lys Asn Met Gly Gln Asn Gln Ile 705 710 715 720

Asn Asn Ile Ala Asp Arg Gly His Ser Tyr Ala Ala Ala Val Ser Ser 725 730 735

Ser Lys Leu Thr Pro Ser Lys Tyr Ile Ser Asp Ile Val Ser Gly Leu 740 745 750

Ser Gln Val Gln Phe Val Met Glu Leu Asn Ser Lys Leu Glu Ser Glu 755 760 765

Gly Lys Glu Tyr Leu Ala Lys Glu Ile Ile Pro Ile Leu Gln Glu Ile 770 775 780

Gln Lys Tyr Val Leu Gln Gly Glu Phe Arg Tyr Arg Leu Val Gly Asn 785 790 795 800

Gln Glu Ile Ile Val Glu Asn Glu Lys Leu Ile Glu Lys Phe Asp Lys 805 810 815 Asp Ile Ser Ser Asn Arg Pro Thr Leu Ser Leu Thr Val Thr Asp Gly 820 825 830

Leu Ser Ala Leu Leu Asn Ser Phe Asn Tyr Asn His Thr Ser Glu Asn 835 840 845

Val Leu Val Asn Leu Pro Phe Gln Val Gly Tyr Ser Ser Leu Gly Lys 850 855 860

Ile Gly Ser Ser Tyr Ser Ser Lys Asp Gly Ala Ser Leu Gln Ile Leu 865 870 875 880

Ser Gln Leu Tyr Ser Phe Lys Asn Leu His Ser Lys Ile Arg Glu Ser 885 890 895

Asn Gly Ala Tyr Gly Gly Gly Leu Thr Tyr Asp Gly Leu Asn Gly Thr 900 905 910

Leu Asn Phe Tyr Ser Tyr Arg Asp Pro Asn Pro Val Lys Ser Ile Gln 915 920 925

Thr Phe Arg Asp Ser Leu Ser Tyr Gly Leu Asp Ala Asn Trp Asn Asp 930 935 940

Lys Asp Leu Gln Glu Ala Lys Leu Arg Val Phe Gln Ser Val Asp Ala 945 950 955 960

Pro Ile Asn Ile Ser Ser Gln Gly Ala Ser Ala Phe Phe Glu Asn Ile 965 970 975

Asp Asp Tyr Leu Arg Gln Glu Arg Arg Glu Asn Phe Leu Gly Thr Thr
980 985 990

Leu Lys Asp Leu Arg Asp Val Thr Glu Lys Tyr Leu Val Asp Asn Gln 995 1000 1005

Asn Asn Leu Val Thr Val Ile Gly Asp Asn Glu Ile Leu Asn Val Asp 1010 1015 1020

Asn Lys Trp Gln Ile Arg Asn Phe Gln Val 1025 1030

<210> 329

<211> 1366

<212> DNA

<213> Candida albicans

```
<400> 329
ttaagaacta gcagatgtaa aatgttttat gtcattatat aatttgttaa tacatgtata 60
tagatttttt aatgaatgta ttccctaaat agaacagaat tatgatgctg ttacagcaaa 120
aactggtttg agtattggag aatgttataa actgaaattt gatttacaac caaaacccgt 180
gtcacgtgta agtaattagg gcttttaggg ctttctatat acaggcacca gaatttttt 240
ttatgggcga tagaaataat gtatgcgcgc gattcttcct gctagaggtt tttctttttg 300
tatgtctgta acagtgcgac tcacatagtt aagtaatttt aaagccagag attgtgtaca 360
gtcgcacgcc ccctaatcac atagttaatt ctcaaacttt ctcttttct ctctttgctt 420
ttctcctaag gaaaaaaaa atttattca tttgttgaaa atttttgtat agttcagttt 480
aacaacacca gtaatcaaac atgccattag ttgtccaaga acaaggttca ttccaacaca 540
<del>ttttacggta tgtgagtgta ttataagatt atttgaatga gatagaaaag ggttaaggaa 600</del>
ggaaagaaat tatccaacta ctgttttggg aaagattaat cagaaatttg aatgcaatga 660
aaaaactttt tcaagagatg gatatggaac aagattcgat tgttatcaac aaatataaag 720
aaaagcaatg atttactact gtttgggaaa gattttggat atgtgtatca acaatactac 780
tatttacttt ggagtaatat gttttcaata ttatcagaga cttatcaaaa tttatgagtt 840
tattattatt gaagttcaat accatcaagc ctttttttta accatcactc tattcaacaa 900
tcatttaaat attaaagtct ttttttccc tgtttaaagt attcgattta ctaacattat 960
gaattattat tatagtttgt taaacaccaa cattgatggt agaatcaaga tcatgtacgc 1020
cttgaccaag atcagaggtg tcggtagaag atatgccaac ttggtttgta aaaaagccga 1080
tgttgaatta accaaaagag ctggtgaatt gacccaagaa gaattggaaa gaattgtcac 1140
cattatgcaa aacccaacca actataaaat cccagcttgg ttcttgaaca gacaaaaaga 1200
tcaagttgat ggtaaagatt accatgtttt agctaacaac ttggaatcta aattgagaga 1260
tgatttggaa agattgaaga aaatcagatc tcacagaggt attagacact tctggggatt 1320
gaaagttaga ggtcaacata ctaaaactac ttctcgtggt cgttaa
                                                                   1366
<210> 330
<211> 145
<212> PRT
<213> Candida albicans
<400> 330
Met Pro Leu Val Val Gln Glu Gln Gly Ser Phe Gln His Ile Leu Arg
  1
                  5
                                     10
Leu Leu Asn Thr Asn Ile Asp Gly Arg Ile Lys Ile Met Tyr Ala Leu
             20
                                 25
```

Thr Lys Ile Arg Gly Val Gly Arg Arg Tyr Ala Asn Leu Val Cys Lys 35 40 45

Lys Ala Asp Val Glu Leu Thr Lys Arg Ala Gly Glu Leu Thr Gln Glu 50 55 60

Glu Leu Glu Arg Ile Val Thr Ile Met Gln Asn Pro Thr Asn Tyr Lys
65 70 75 80

SEQL

```
Ile Pro Ala Trp Phe Leu Asn Arg Gln Lys Asp Gln Val Asp Gly Lys
85 90 95
```

Asp Tyr His Val Leu Ala Asn Asn Leu Glu S r Lys Leu Arg Asp Asp 100 105 110

Leu Glu Arg Leu Lys Lys Ile Arg Ser His Arg Gly Ile Arg His Phe 115 120 125

Trp Gly Leu Lys Val Arg Gly Gln His Thr Lys Thr Thr Ser Arg Gly 130 135 140

Arg 145

<210> 331 <211> 1327 <212> DNA <213> Candida albicans

<400> 331

atagttatta caatataata aagcaaataa ataaaagaaa tgataaagaa ccatattaac 60 aaagtttgaa cgtgctgtaa gtaaatcttg tatacgagaa tattcatctt ggaatatttc 120 gccctaatag tatatgcagt cgcatacatt aattggtcat ctcataagta aattatataa 240 tgtatagtgt gcgcgcactg taatttttca cactaagtta aggagagtgc gaaaaattta 300 gtactaacaa aattgtattg tgttgtgtgt ggctattggg cagagcgaaa atttcacccc 360 tactgagagg aaactgtgag ggagagagat acacacaaac tctgtgcgca agaagaaaga 420 caaaaaattt ttttgaaaaa aaaaaaagag acaaccaacc ttttattgta ttaacatttt 480 aagatatcaa gaaactaaaa atggctaagt tcatcaaatc tggtaaagtt ggtatgtaaa 540 aatgagtacc tatatgggta gacaatgata gatttattgg ccaactatga atgggataga 600 tatatatgat atgggagaga attttgaagg attgaaatat tttatataaa agagagaagc 660 aaagagaaat aatgaacaat tgaaaagaag atacaaatac tgttaattaa actagaatat 720 gccattgaag atttaaatga ttaccatcgt aataaattta ttggaatttg ggagattctt 780 ttggagttaa gaaaaaagca agtgattgat caatcactat ggagccgtca caagacatca 840 acgaatccca caacattttt atttattcat atccatcata tggcagtaac aatcatgaat 900 ctacaaatta catttaaaaa aagcattata ctaactaaat tttttagcta ttgttgtaag 960 aggtegttae getggtaaaa aagtagteat tgtgaaacea catgatgaag gtaccaaate 1020 tcacccattc ccacatgcca ttgttgctgg tattgaaaga gctccattga aggttaccaa 1080 gaagatggat gctaaaaaag ttaccaaaag aactaaagtc aagccatttg ttaaattagt 1140 aaactacaac catttaatgc caactagata ctcattagat gttgaatcat tcaaatctgc 1200 tgtcacttct gaagctttag aagaaccatc tcaaagagaa gaagctaaaa aagttgtcaa 1260 gaaggctttt gaagaaaaac atcaagctgg taagaacaaa tggttcttcc aaaaattaca 1320 cttttaa 1327

```
<210> 332
```

<211> 136

<212> PRT

<213> Candida albicans

<400> 332

Met Ala Lys Phe Ile Lys Ser Gly Lys Val Ala Ile Val Val Arg Gly
1 5 10 15

Arg Tyr Ala Gly Lys Lys Val Val Ile Val Lys Pro His Asp Glu Gly 20 25 30

Thr Lys Ser His Pro Phe Pro His Ala Ile Val Ala Gly Ile Glu Arg
35 40 45

Ala Pro Leu Lys Val Thr Lys Lys Met Asp Ala Lys Lys Val Thr Lys 50 55 60

Arg Thr Lys Val Lys Pro Phe Val Lys Leu Val Asn Tyr Asn His Leu 65 70 75 80

Met Pro Thr Arg Tyr Ser Leu Asp Val Glu Ser Phe Lys Ser Ala Val 85 90 95

Thr Ser Glu Ala Leu Glu Glu Pro Ser Gln Arg Glu Glu Ala Lys Lys
100 105 110

Val Val Lys Lys Ala Phe Glu Glu Lys His Gln Ala Gly Lys Asn Lys
115 120 125

Trp Phe Phe Gln Lys Leu His Phe 130 135

<210> 333

<211> 1157

<212> DNA

<213> Candida albicans

<400> 333

ttccagtcat tccgaaatgc ataagcaaga cgttagttgt attgtggttg tttagtctaa 60 tccaaaaata cttgccatga cgacccactg aagtaacaac tgcatttgtt aaggcttttc 120 tcatcttatt caattcttt tcagcatcat tcgtatcttt aagcacaggg aatagcaatg 180 gatcatgtaa caaattcact ttcgtgattc gaaagcccaa aatgtttcgt tttaacaacg 240 cacacacatg tgaaacctaa accgagttag tcgtcaagaa aataattcag tgttaaagtc 300 tgtaccttaa gcgtcaaacg tacttctgca acctctggca ttgagtgtaa tttaaatatt 360 catgataatc tggaactaca agctacgaaa aaaaaagaaa aagaaaaata gtacgagttc 420

```
ttggtgagat aaataatgga gaacacttt ttttttctt ttggaggttt tagaaggcaa 480
aaccaaatct agacaaggag atgaacagat tattcggaac gaaaagcact gcgcccaaac 540
catctttgaa tgatgcaatc aagggaatcg atgaaagagt ggggtctttg gatgttaaat 600
taagcaagat caactcggaa ttatccacat accaacaaa gataagcaga atgaaggagg 660
gacccgggaa gtcggcacta aaacaaaaag caatcaagtt actaagacag aggaagcaga 720
tagaagctca aaaggatcag ttagagaatc aatcttggaa tatgacacaa gcttccatga 780
caacagataa cttacaaaat accatggtca caataaatgc aatgaaaaca gccaataaac 840
tgttgaaaca aacttatgga aaaattaata tcgatgaatt ggaagatctt caagatgaaa 900
tgttggattt gattgataaa tcaaatgaac tacaggaggc acttctgacg agctatgatg 960
tagattttga aaatgaaatg gcagaggtg ggataggtgc acctagttac ttaaatgata 1020
ttgattttga aaatgaaatg gcagagagtg ggataggtgc acctagttac ttaaatgata 1080
cagaacctac agcagcagat aaattgccta catttattga cgaacaacca gaagaagctc 1140
aaaaaaatcgc aaactag
```

<210> 334

<211> 218

<212> PRT

<213> Candida albicans

<400> 334

Met Asn Arg Leu Phe Gly Thr Lys Ser Thr Ala Pro Lys Pro Ser Leu
1 5 10 15

Asn Asp Ala Ile Lys Gly Ile Asp Glu Arg Val Gly Ser Leu Asp Val 20 25 30

Lys Leu Ser Lys Ile Asn Ser Glu Leu Ser Thr Tyr Gln Gln Lys Ile 35 40 45

Ser Arg Met Arg Asp Gly Pro Gly Lys Ser Ala Leu Lys Gln Lys Ala 50 55 60

Ile Lys Leu Leu Arg Gln Arg Lys Gln Ile Glu Ala Gln Lys Asp Gln 65 70 75 80

Leu Glu Asn Gln Ser Trp Asn Met Thr Gln Ala Ser Met Thr Thr Asp 85 90 95

Asn Leu Gln Asn Thr Met Val Thr Ile Asn Ala Met Lys Thr Ala Asn 100 105 110

Lys Ser Leu Lys Gln Thr Tyr Gly Lys Ile Asn Ile Asp Glu Leu Glu 115 120 125

Asp Leu Gln Asp Glu Met Leu Asp Leu Ile Asp Lys Ser Asn Glu Leu 130 135 140

Gln Glu Ala Leu Ser Thr Ser Tyr Asp Val Pro Asp Asp Ile Ser Glu 145 150 155 160

Ser Glu Leu Asp Ala Glu Leu Glu Ala Leu Gly Glu Glu Ile Asp Phe 165 170 175

Glu Asn Glu Met Ala Glu Ser Gly Ile Gly Ala Pro Ser Tyr Leu Asn 180 185 190

Asp Thr Glu Pro Thr Ala Ala Asp Lys Leu Pro Thr Phe Ile Asp Glu 195 200 205

Gln Pro Glu Glu Ala Gln Lys Ile Ala Asn 210 215

<210> 335 <211> 4550 <212> DNA

<213> Candida albicans

<400> 335 gaatgagatt ttttttttta ctaagggtgc actactacta gttatttgtt ttgttgttga 60 cgatcattaa agaaaaaatt tacagtatac acaaaacact ttacttctgc tgtttttttt 120 ttattttagt ttttggtttc ataaattatt aaaagaaagc aaataattat tgaaataaat 180 tttatatttt tggtttttt ttctttggtt tctttgaatt ttgcaaacca atccaaattt 240 tttttgaaaa tttttccttc ttcttcattt gttgactttt gaaagtttta ttcatccata 300 ttcttcaagt taaaatagtg ttcaatatct gtccaaccaa gaagaaaacc aataaacaac 360 atcaattcca actttgttta catcaaacag aaacaaaaaa cagataattt atagacacct 420 tcattcattc gttattttt tccaagtact acacgtcttt atttttaata catttatcat 480 . aaacaaaatt tatatcaata atgccacatt caagacaacc ttcgatatcg tcatcgatta 540 tgtcccaatc aaatcataat catccacaga agatcggacc ttggaaatta ggtaaaactt 600 taggaagagg tgccaccggt agagtcttat tagctactca tcaaactact ggtcaaaaag 660 ctgccgttaa agtagtttcc aaatccgaat tacaagacga agaaaccgag aaaaatggag 720 atggattacc atatggtata gaaagagaaa ttattataat gaaattatta actcatccta 780 atgittigag attatatgat gtatgggaaa catctaaggc tctttatctt gttctagaat 840 acgttgaagg tggagaatta tttgatttat tggtggaacg tggtccatta cctgaagtcg 900 aggetateaa gtattteegt caaateatat tgggtaegge ttattgteat getettggta 960 tatgtcacag agatttaaaa ccagagaatt tgttactaga ttctcaattg aatgtaaaat 1020. tagccgattt tggaatggca gctttagaaa gtaacggcaa attattagag acttcttgtg 1080 gttcacctca ttatgctgct ccagaaatcg ttagtggatt gaaatatcat ggagctgctt 1140 ctgatgtttg gtcttgtggg gttatattat ttgccttgtt gacaggtaga ttaccctttg 1200 atgatgaaaa tattagaaat ttacttctta aagtccaggc tggtaacttt gaaatgcccg 1260 ttgatgaagt tagcagagaa gctagagatt taattgctag aatgttagag gttgatccta 1320 tgagaagaat atctactgaa aaaatcttaa gacacccatt gttaaccaaa tacccaatgt 1380 caaacgaaga tttaatcagt gaaaaatcat taccacatcc acatactggt tacaaatctt 1440 tagggtcagt tagaaacatt gataaacaga ttttatcaaa cttgacaatt ttatggaatg 1500

```
atagacccga agaggaaatt gttgattgtc ttttgaaaga tggatccaat ccagaaaaaa 1560
cattctatgc attattgatg agatacaagc ataatcaaga cgataacact aataacaatt 1620
caccaaagaa atcaacgagt ttcaataata aagtggtacg cagtgggtcc aaatacagtc 1680
ttaatggaac ccctagaaga aaaagagcca gtcacataag tgtgtcaaga ccaacatctt 1740
tccaatacaa gtctaatcct ggcgctggtg caacagcaaa tagaaactcc gttgccagac 1800
attotgtggc ttcctcggcc aacaattoto otogtaaato accatacaag tcaccataca 1860
gatcacctta tagatcacca tataaatcgc cttctaagag atattcatat aatcaatccc 1920
caactaaatc tccttacgga agaagatcaa attcacaaag acaatttgaa aatgaaccat 1980
taaaggcaaa gccaagaaat atttacaatg agattgttga tgcacaaagc aacttttctc 2040
tacctccatc gcttccacct tccttacctt caaaagattc tcgttatatg atcgatgaac 2100
ccaatcaacc ccagttgcaa caacctgctt taagtcaagt ccctgaaaat cctattgttg 2160
atgaatcccc tgatttaatg cagtcagcaa aaatttctag tggaaagaga aattcaataa 2220
taggaaagaa caacaacaac agcaacagca acaagagaat gtctaagaga aaatcaattc 2280
gtgcatctat gaccacggga ttgaaaagaa attccataac catgaaattg ttatctactt 2340
atgctaaatt atcaggtgat gacgactggg aatacatgga taaacaaaca aaaagaacat 2400
eggetaettt tgeageattg tgtgacaaaa tatttaatca agaagaetat gaegaagaag 2460
acgaacaatt agttgatcct gaagaaaagg aagccaagga atatgaaagg ttaatggaat 2520
tggaaagaaa gaaacatgaa gctgagttga aagctagaag agaattagaa aagaagaaaa 2580
gaagacaaaa gagacgttcc attttgagtt ctaagaaatt aagtattatt gtcaagaatg 2640
atgctgatcc aaataatagt gaacaagagt tggtcgatga aggtataaaa caaccaaaac 2700
gtcaatccaa aaacttgacc gctttaagag cattatctga aggaaatcat gcatctgaag 2760
aattgacatt ggaagacgtt gagaatttga agagacgatc agcatcacag ccggttccaa 2820
aaagaagaca aactccggtt ttgacaagaa gacctgtatc aagattagat ccattatggc 2880
aagcacacga gaatgaacag ttagatagag caaaagatgc cttggaacaa gaatggaggg 2940
attcacaaaa gagaagttct actgttagtc gtaaaaaagt caacagagag tcgatgatat 3000
cagttatgga tgatattgtt gaagaggacc aaggccgtgt caacaggaga tcaacacgaa 3060
acacttacta tgaaagggaa agagactatg aattaccaga accaactgtg gaagattcca 3120
acttgactga tgactacatg acagaaatca gaaaatcaag acttttgaac agtcagttaa 3180
atgttaggga tccacttaat gaaaaaagaa aatctgaacc caagactctt attagcaatg 3240
ttcaaatacc gagtgttact agaaaatcaa gaaatttcac tacttccaac aaaaggttgt 3300
cggtattgtc tatgtattca acaaaggaat cataccgtga tttgaattct ataattaact 3360
caccagatga aaaccccgaa cagcatcaaa acatgaacaa gccagcgtta cgaaccagta 3420
ttgctgatcg tttggataaa gctggattgg ctgaaccaga atatgaaact gagactgatg 3480
gtgaagataa agtgtctgtt attgatttgg atgatcattt agctgataga aggacttcct 3540
attatgatgg atctggaaag agagcatcta gagcttcaac aactaaacgt tacaatgttc 3600
attccagttc agaaaaaaga ccaaaatcca aagttcctga tttgccaaag aatgattatg 3660
atgacacatt tgtcagtaat agtgatgaag ttcataagcg tcagtataaa tcgatggttt 3720
ctgatgagtc tagtgcatct gatgatgtat ttgataagat taaattacca gatggtaaat 3780
caactaaatc ttccattgat gaattggcta acggcacgtc tacaagtggt catagaaaac 3840
caaagataag acatteteaa eegggeeeag aaatgttgat teeteatttg aatggaggta 3900
ttgagtcgtc tcaaccaatg tctaaagttc gtggtaacaa ttcaagtggt catgatgata 3960
gtgttccacc accaccgcca gctcacaagg tgaataaaaa accattggat gataagacga 4020
atttccctcc accagaagtg gatccaaaaa gaaaaggttc attttttaga aaactttctt 4080
ggggatccaa aaaaaccatt gaaaataata caaacgccgc cactaatacc acgactcaac 4140
aacaattacc aagtcctgct gaatcaaaag aggagaaacc aaaaagttca tttttcagat 4200
ggttttcgtc atctaatact ccatctgctg ctgaaattag aaaattcaac accattttac 4260
ctaaacatga aatgtctact gctttatttg ctttattgaa ttcttggtct aattttggtt 4320
tgaaagattt acggaatgat caagttggat attatattac tggtgctatt tctaaacata 4380
```

SEQL

attetttaa titaaagagt tigaaatta gaattaagat taateaaga gattitaate 4440 aaaaateaga aattittigt gitagagtga aaggatetaa agitaeaact gataettat 4500 titigigaaat tigaaaaggte tiaeteaaag aaggiggtit agataaataa 4550

<210> 336

<211> 1349

<212> PRT

<213> Candida albicans

<400> 336

Met Pro His Ser Arg Gln Pro Ser Ile Ser Ser Ser Ile Met Ser Gln

1

5

10

15

Ser Asn His Asn His Pro Gln Lys Ile Gly Pro Trp Lys Leu Gly Lys
20 25 30

Thr Leu Gly Arg Gly Ala Thr Gly Arg Val Leu Leu Ala Thr His Gln
35 40 45

Thr Thr Gly Gln Lys Ala Ala Val Lys Val Val Ser Lys Ser Glu Leu 50 55 60

Gln Asp Glu Glu Thr Glu Lys Asn Gly Asp Gly Leu Pro Tyr Gly Ile
65 70 75 80

Glu Arg Glu Ile Ile Ile Met Lys Leu Leu Thr His Pro Asn Val Leu 85 90 95

Arg Leu Tyr Asp Val Trp Glu Thr Ser Lys Ala Leu Tyr Leu Val Leu
100 105 110

Glu Tyr Val Glu Gly Glu Leu Phe Asp Leu Leu Val Glu Arg Gly
115 120 125

Pro Leu Pro Glu Val Glu Ala Ile Lys Tyr Phe Arg Gln Ile Ile Leu 130 135 140

Gly Thr Ala Tyr Cys His Ala Leu Gly Ile Cys His Arg Asp Leu Lys 145 150 155 160

Pro Glu Asn Leu Leu Asp Ser Gln Leu Asn Val Lys Leu Ala Asp 165 170 175

Phe Gly Met Ala Ala Leu Glu Ser Asn Gly Lys Leu Leu Glu Thr Ser 180 185 190

Cys Gly Ser Pro His Tyr Ala Ala Pro Glu Ile Val Ser Gly Leu Lys

SEQL

195

205

Tyr His Gly Ala Ala Ser Asp Val Trp Ser Cys Gly Val Ile Leu Phe 210 215 220

200

Ala Leu Leu Thr Gly Arg Leu Pro Phe Asp Asp Glu Asn Ile Arg Asn 225 230 235 240

Leu Leu Lys Val Gln Ala Gly Asn Phe Glu Met Pro Val Asp Glu 245 250 255

Val Ser Arg Glu Ala Arg Asp Leu Ile Ala Arg Met Leu Glu Val Asp
260 265 270

Pro Met Arg Arg Ile Ser Thr Glu Lys Ile Leu Arg His Pro Leu Leu 275 280 285

Thr Lys Tyr Pro Met Ser Asn Glu Asp Leu Ile Ser Glu Lys Ser Leu 290 295 300

Pro His Pro His Thr Gly Tyr Lys Ser Leu Gly Ser Val Arg Asn Ile 305 310 315 320

Asp Lys Gln Ile Leu Ser Asn Leu Thr Ile Leu Trp Asn Asp Arg Pro 325 330 335

Glu Glu Glu Ile Val Asp Cys Leu Leu Lys Asp Gly Ser Asn Pro Glu 340 345 350

Lys Thr Phe Tyr Ala Leu Leu Met Arg Tyr Lys His Asn Gln Asp Asp 355 360 365

Asn Thr Asn Asn Asn Ser Pro Lys Lys Ser Thr Ser Phe Asn Asn Lys 370 375 380

Val Val Arg Ser Gly Ser Lys Tyr Ser Leu Asn Gly Thr Pro Arg Arg 385 390 395 400

Lys Arg Ala Ser His Ile Ser Val Ser Arg Pro Thr Ser Phe Gln Tyr
405 410 415

Lys Ser Asn Pro Gly Ala Gly Ala Thr Ala Asn Arg Asn Ser Val Ala 420 425 430

Arg His Ser Val Ala Ser Ser Ala Asn Asn Ser Pro Arg Lys Ser Pro 435 440 445

Tyr Lys Ser Pro Tyr Arg Ser Pro Tyr Arg Ser Pro Tyr Lys Ser Pro

Ser Lys Arg Tyr Ser Tyr Asn Gln Ser Pro Thr Lys Ser Pro Tyr Gly Arg Arg Ser Asn Ser Gln Arg Gln Phe Glu Asn Glu Pro Leu Lys Ala Lys Pro Arg Asn Ile Tyr Asn Glu Ile Val Asp Ala Gln Ser Asn Phe Ser Leu Pro Pro Ser Leu Pro Pro Ser Leu Pro Ser Lys Asp Ser Arg Tyr Met Ile Asp Glu Pro Asn Gln Pro Gln Leu Gln Gln Pro Ala Leu Ser Gln Val Pro Glu Asn Pro Ile Val Asp Glu Ser Pro Asp Leu Met Gln Ser Ala Lys Ile Ser Ser Gly Lys Arg Asn Ser Ile Ile Gly Lys Asn Asn Asn Ser Asn Ser Asn Lys Arg Met Ser Lys Arg Lys Ser Ile Arg Ala Ser Met Thr Thr Gly Leu Lys Arg Asn Ser Ile Thr Met Lys Leu Leu Ser Thr Tyr Ala Lys Leu Ser Gly Asp Asp Trp Glu Tyr Met Asp Lys Gln Thr Lys Arg Thr Ser Ala Thr Phe Ala Ala Leu Cys Asp Lys Ile Phe Asn Gln Glu Asp Tyr Asp Glu Glu Asp Glu Gln 

Leu Val Asp Pro Glu Glu Lys Glu Ala Lys Glu Tyr Glu Arg Leu Met 

Glu Leu Glu Arg Lys Lys His Glu Ala Glu Leu Lys Ala Arg Arg Glu 

Leu Glu Lys Lys Lys Arg Arg Gln Lys Arg Arg Ser Ile Leu Ser Ser 

Lys Lys Leu Ser Ile Ile Val Lys Asn Asp Ala Asp Pro Asn Asn Ser

Glu Gln Glu Leu Val Asp Glu Gly Ile Lys Gln Pro Lys Arg Gln Ser Lys Asn Leu Thr Ala Leu Arg Ala Leu Ser Glu Gly Asn His Ala Ser Glu Glu Leu Thr Leu Glu Asp Val Glu Asn Leu Lys Arg Arg Ser Ala Ser Gln Pro Val Pro Lys Arg Arg Gln Thr Pro Val Leu Thr Arg Arg Pro Val Ser Arg Leu Asp Pro Leu Trp Gln Ala His Glu Asn Glu Gln Leu Asp Arg Ala Lys Asp Ala Leu Glu Glu Trp Arg Asp Ser Gln Lys Arg Ser Ser Thr Val Ser Arg Lys Lys Val Asn Arg Glu Ser Met Ile Ser Val Met Asp Asp Ile Val Glu Glu Asp Gln Gly Arg Val Asn Arg Arg Ser Thr Arg Asn Thr Tyr Tyr Glu Arg Glu Arg Asp Tyr Glu Leu Pro Glu Pro Thr Val Glu Asp Ser Asn Leu Thr Asp Asp Tyr Met Thr Glu Ile Arg Lys Ser Arg Leu Leu Asn Ser Gln Leu Asn Val Arg Asp Pro Leu Asn Glu Lys Arg Lys Ser Glu Pro Lys Thr Leu Ile Ser 

Asn Val Gln Ile Pro Ser Val Thr Arg Lys Ser Arg Asn Phe Thr Thr

Ser Asn Lys Arg Leu Ser Val Leu Ser Met Tyr Ser Thr Lys Glu Ser 

Tyr Arg Asp Leu Asn Ser Ile Ile Asn Ser Pro Asp Glu Asn Pro Glu 

Gln His Gln Asn Met Asn Lys Pro Ala Leu Arg Thr Ser Ile Ala Asp

955

970

975

Arg Leu Asp Lys Ala Gly Leu Ala Glu Pro Glu Tyr Glu Thr Glu Thr 980 985 990

Asp Gly Glu Asp Lys Val Ser Val Ile Asp Leu Asp Asp His Leu Ala 995 1000 1005

Asp Arg Arg Thr Ser Tyr Tyr Asp Gly Ser Gly Lys Arg Ala Ser Arg 1010 1015 1020

Ala Ser Thr Thr Lys Arg Tyr Asn Val His Ser Ser Ser Glu Lys Arg 1025 1030 1035 1040

Pro Lys Ser Lys Val Pro Asp Leu Pro Lys Asn Asp Tyr Asp Asp Thr 1045 1050 1055

Phe Val Ser Asn Ser Asp Glu Val His Lys Arg Gln Tyr Lys Ser Met 1060 1065 1070

Val Ser Asp Glu Ser Ser Ala Ser Asp Asp Val Phe Asp Lys Ile Lys 1075 1080 1085

Leu Pro Asp Gly Lys Ser Thr Lys Ser Ser Ile Asp Glu Leu Ala Asn 1090 1095 1100

Gly Thr Ser Thr Ser Gly His Arg Lys Pro Lys Ile Arg His Ser Gln 1105 1110 1115 1120

Pro Gly Pro Glu Met Leu Ile Pro His Leu Asn Gly Gly Ile Glu Ser 1125 1130 1135

Ser Gln Pro Met Ser Lys Val Arg Gly Asn Asn Ser Ser Gly His Asp 1140 1145 1150

Asp Ser Val Pro Pro Pro Pro Pro Ala His Lys Val Asn Lys Lys Pro 1155 1160 1165

Leu Asp Asp Lys Thr Asn Phe Pro Pro Pro Glu Val Asp Pro Lys Arg 1170 1175 1180

Lys Gly Ser Phe Phe Arg Lys Leu Ser Trp Gly Ser Lys Lys Thr Ile 1185 1190 1195 1200

Glu Asn Asn Thr Asn Ala Ala Thr Asn Thr Thr Thr Gln Gln Gln Leu 1205 1210 1215

Pro Ser Pro Ala Glu Ser Lys Glu Glu Lys Pro Lys Ser Ser Phe Phe

1220

1225

1230

Arg Trp Phe Ser Ser Ser Asn Thr Pro Ser Ala Ala Glu Ile Arg Lys 1235 1240 1245

Phe Asn Thr Ile Leu Pro Lys His Glu Met Ser Thr Ala Leu Phe Ala 1250 1255 1260

Leu Leu Asn Ser Trp Ser Asn Phe Gly Leu Lys Asp Leu Arg Asn Asp 1265 1270 1275 1280

Gln Val Gly Tyr Tyr Ile Thr Gly Ala Ile Ser Lys His Asn Ser Phe 1285 1290 1295

Asn Leu Lys Ser Cys Lys Phe Arg Ile Lys Ile Asn Gln Arg Asp Phe 1300 1305 1310

Asn Gln Lys Ser Glu Ile Val Cys Val Arg Val Lys Gly Ser Lys Val 1315 1320 1325

Thr Thr Asp Thr Leu Phe Cys Glu Ile Glu Lys Val Leu Leu Lys Glu 1330 1335 1340

Gly Gly Leu Asp Lys 1345

<210> 337

<211> 1121

<212> DNA

<213> Candida albicans

<400> 337

acaatactag gcactgttga gtgagtgagc attittictg titetcactc agttaacaaa 60 ataaaaaaaa titecataat titagaagtit cattacagt cititticaa titaacagtga 120 tacaaagagtg tatgtaaaga caacatgtac tagcaactat aatatgatit accaatgati 180 gggatcacaa taaatgtgit aatatgaatg agagaaggat agtgaataag agattacgaa 240 agaatagati caacaagtic agaatggtat acaactaaaa tggaattati titcaaatatg 300 caactatcat tatgactact acgacaacaa tittaatcga gagaaggatca titagatcaag 360 agitgggaaa ctaataccaa ggaaatatca titaagaatta atagcitigc aaaaattggit 420 titactcata titattigiti tagitggaaa gcgattacat catggaacaa agittactaa 480 caacattgit ataggtaaaa atgggtatti citagagatic acgicacaaa agatccgcca 540 ciggigccaa aagagcccaa ticagaaaga agagaaagti tigaattaggi agacaaccag 600 ccaacaccaa gattggicca aaaagaattc actitgicag aaccagaggi ggtaaccaaa 660 aattcagagc titigagagti gaaaccggta actitcitig gggticcgaa ggtgittcca 720 gaaaaaccag aattgctggi gtcgittacc atcatcaa taacgaattg caatggtacg 840 acaccitigac caaatctgct gttgitcaaa tigatgctac tccattcag caatggtacg 840

aaaaccacta cggtgctact ttaggtaaaa agaagggtgg tgctcatgct gctcacgctg 900 ctgaagttgc cgatgccaag agatcaagaa aagtcgaaag aaaattggct gctagatctg 960 gtgctgctgc cattgaatcc gctgttgact ctcaattcgg ttctggtaga ttatacgctg 1020 tcatttcttc aagaccaggt caatctggta gatgtgatgg ttacatcttg gaaggtgaag 1080 aattagcctt ctacttgaga agattaactg ctaagaaata a 1121

<210> 338

<211> 206

<212> PRT

<213> Candida albicans

<400> 338

Met Gly Ile Ser Arg Asp Ser Arg His Lys Arg Ser Ala Thr Gly Ala
1 5 10 15

Lys Arg Ala Gln Phe Arg Lys Lys Arg Lys Phe Glu Leu Gly Arg Gln 20 25 30

Pro Ala Asn Thr Lys Ile Gly Pro Lys Arg Ile His Ser Val Arg Thr
35 40 45

Arg Gly Gly Asn Gln Lys Phe Arg Ala Leu Arg Val Glu Thr Gly Asn 50 55 60

Phe Ser Trp Gly Ser Glu Gly Val Ser Arg Lys Thr Arg Ile Ala Gly 65 70 75 80

Val Val Tyr His Pro Ser Asn Asn Glu Leu Val Arg Thr Asn Thr Leu 85 90 95

Thr Lys Ser Ala Val Val Gln Ile Asp Ala Thr Pro Phe Arg Gln Trp
100 105 110

Tyr Glu Asn His Tyr Gly Ala Thr Leu Gly Lys Lys Lys Gly Gly Ala 115 120 125

His Ala Ala His Ala Ala Glu Val Ala Asp Ala Lys Arg Ser Arg Lys
130 135 140

Val Glu Arg Lys Leu Ala Ala Arg Ser Gly Ala Ala Ala Ile Glu Ser

Ala Val Asp Ser Gln Phe Gly Ser Gly Arg Leu Tyr Ala Val Ile Ser 165 170 175

Ser Arg Pro Gly Gln Ser Gly Arg Cys Asp Gly Tyr Ile Leu Glu Gly 180 185 190

Glu Glu Leu Ala Phe Tyr Leu Arg Arg Leu Thr Ala Lys Lys 195 200 205

<210> 339

<211> 819

<212> DNA

<213> Candida albicans

<400> 339

ttctcctgtg aaaagtttcg agatgtaacg tttcgcagta atagagagcc agaatccatt 60
tttgtgtact acagacaaat tcagaagttt caactgctgc atatcgcctt aaatgactgt 120
agcattcgtc caaattgaga ccctcaatta cattttgtca aaaaaattgg tccctagtgt 180
tgctatcgat aacgaaggtg aaggcagttt agcttggaga catttagaga acttagttac 240
atctcatctt ccgtttcgag aaatcgttga tttaccgtgc agcgcttata ttgattgcta 300
cttgttccca gcaccacagc aatatagcaa tcataaataa attgccccgc ggttgacagt 360
gtatatcttc gaggaatggc aacctttgcc cccctctcga aaaacaatat aaatagagtc 420
aatttctcta gtagaggtaa attctttgaa tcttgtttt tttcgacata caccataaat 480
cccatagaaa actgcaaaat gtctgacgcc ggaagaaaaa acatttctac taaaatcaac 540
gaagctataa cccccgaatc cgaaaagtct accttggaaa agggcaagga acaagtcacc 600
agtacccttg acaaagctgt tggctcaaat gttccagata accaaaaatc tttcactcaa 660
actgttgcag acagcgtgca acaaagaccc ttgctgaaac agctcaagaa tatgtcgagg 780
ttgccaaaac tgaaattgga aaggctgctg aatacgtga

<210> 340

<211> 106

<212> PRT

<213> Candida albicans

<400> 340

Met Ser Asp Ala Gly Arg Lys Asn Ile Ser Thr Lys Ile Asn Glu Ala

1 5 10 15

Ile Thr Pro Glu Ser Glu Lys Ser Thr Leu Glu Lys Gly Lys Glu Gln

20 25 30

Val Thr Ser Thr Leu Asp Lys Ala Val Gly Ser Asn Val Pro Asp Asn 35 40 45

Gln Lys Ser Phe Thr Gln Thr Val Ala Asp Ser Val Gln Gln Gly Ser 50 55 60

Asp Asn Ala Lys Ala Asp Leu Lys Lys Gln Ser Glu Gln Ala Glu Gly 65 70 75 80



Glu Gln Arg Pro Leu Ser Lys Gln Leu Lys Asn Met Ser Arg Leu Pro 85 90 95

Lys Ser Lys Leu Glu Arg Ser Ser Asn Thr 100 105

<210> 341

<211> 884

<212> DNA

<213> Candida albicans

## <400> 341

ccttctcctg tgaaaagttt cgagatgtaa cgtttcgcag taatagagag ccagaatcca 60 tttttgtgta ctacagacaa attcagaagt ttcaactgct gcatatcgcc ttaaatgact 120 gtagcattcg tccaaattga gaccctcaat tacattttgt caaaaaaaatt ggtccctagt 180 gttgctatcg ataacgaagg tgaaggcagt ttagcttgga ggcatttaga gaacttagtt 240 acateteate tteegttteg agaaategtt gatttacegt geagegetta tattgattge 300 tacttgttcc cagcaccaca gcaatatagc aatcataaat aaattgcccc gcggttgaca 360 gtgtatatct tcgaggaatg gcaacctttg cccccctctc gaaaaacaat ataaatagag 420 tcaatttctc tagtagaggt aaattctttg aatcttgttt tttttcgaca aacaccataa 480 atcccataga aaactgcaaa atgtctgacg ccggaagaaa aaacatttct actaaaatca 540 acgaagctat aacccccgaa tccgaaaagt ctaccttgga aaagggcaag gaacaagtca 600 ccagtaccct tgacaaagct gttggctcaa atgttccaga taaccaaaaa tctttcactc 660 aaactgttgc agacaacgtg caacaaggtt ccgataatgc taaagctgat ttgaagaaac 720 aatccgaaca agcagaggc gaagcaaaga cccttgctga aacagctcaa gaatatgtcg 780 aggttgccaa aactgaaatt ggaaaggctg ctgaatacgt gagtggagtt gtcaccggtg 840 ctaccgaagg tgccaaaacc ggcgctgata gtactaaaaa atag 884

<210> 342

<211> 127

<212> PRT

<213> Candida albicans

<400> 342

Met Ser Asp Ala Gly Arg Lys Asn Ile Ser Thr Lys Ile Asn Glu Ala

1 10 15

Ile Thr Pro Glu Ser Glu Lys Ser Thr Leu Glu Lys Gly Lys Glu Gln
20 25 30

Val Thr Ser Thr Leu Asp Lys Ala Val Gly Ser Asn Val Pro Asp Asn 35 40 45

Gln Lys Ser Phe Thr Gln Thr Val Ala Asp Asn Val Gln Gln Gly Ser

55

60

Asp Asn Ala Lys Ala Asp Leu Lys Lys Gln Ser Glu Gln Ala Glu Gly 65 70 75 80

Glu Ala Lys Thr Leu Ala Glu Thr Ala Gln Glu Tyr Val Glu Val Ala 85 90 95

Lys Thr Glu Ile Gly Lys Ala Ala Glu Tyr Val Ser Gly Val Val Thr
100 105 110

Gly Ala Thr Glu Gly Ala Lys Thr Gly Ala Asp Ser Thr Lys Lys
115 120 125

<210> 343

<211> 1244

<212> DNA

<213> Candida albicans

### <400> 343

gaagggcacc ataatgaaat cgactcactt caggattata atggtatgaa acattgtact 60 tgttattagt gccaggatga ttaggatcat atattggggt gttttctcga gtcttggtat 120 cggttgtaaa cgtatctgtt tcacttatca gtatcgtcat ttatattaac tacttttctc 180 ctatggttat atattggtaa acaaagaaac aaaacaacaa aaaagaagta gtagttttga 240 tgaaagtgag tgcgacataa tgtagaaaaa tgtcgaatgt cttgaacttt acccattgag 360 tagttgttgt agtgtaggag gaagaaaaca acagaaagaa agagagaaag aaaaatttcg 420 ccactacaaa tattcaacaa gtttcatata gtaatataat cccaattgat cattacttta 480 ttccacacaa ttcataaaca atgtccaatt cagcaggttt tgatagacat atcactattt 540 tttctcctga aggtagatta taccaagtag aatatgcttt taaagctatc aattcagcaa 600 atatcaccag tttaggaatc acaggtcaag attctgccgt tattatatca caaaagaaga 660 tcccagataa gttattagat cctaaaaccg tgtcatatat ttttaaaaatc actcctagta 720 taggaatggt tgccactgga tcaattgctg atgctagagc tcaagccatg agagcaagat 780 ctgaagctac agaatttaga tataaatatg gttacgaaat gccggtggaa agtttatcaa 840 gaagaatggc gaatatatct caattgtata ctcaaagagc ttatatgaga ccattgggtg 900 ttgctttaac ttttattcaa gttgattttg ctgatgaagg tagaggtcca caaattttta 960 aatgtgatcc tgctggatat ttcactgggg tgaaagccgt ggccactggt ccaaaacaac 1020 aagaagcaac gacttattta gagaaaaaat tcaaaaaaac cgatgctgtt aaaggagatt 1080 ggcaaaaaac tgttgaattt gcaataattg ccttgagttc tgtgattgga actgaattca 1140 gaaaaaatga tattgaaatt ggtgttgcca ctgaaggaga atttagaatt ttgacaccag 1200 aagaaataga cgaaagattg atttcaatag ctgaacaaga ttag 1244

<210> 344

<211> 247

<212> PRT

<213> Candida albreans

<400> 344

Met Ser Asn Ser Ala Gly Phe Asp Arg His Ile Thr Ile Phe Ser Pro 1 5 10 15

Glu Gly Arg Leu Tyr Gln Val Glu Tyr Ala Phe Lys Ala Ile Asn Ser 20 25 30

Ala Asn Ile Thr Ser Leu Gly Ile Thr Gly Gln Asp Ser Ala Val Ile 35 40 45

Ile Ser Gln Lys Lys Ile Pro Asp Lys Leu Leu Asp Pro Lys Thr Val
50 55 60

Ser Tyr Ile Phe Lys Ile Thr Pro Ser Ile Gly Met Val Ala Thr Gly 65 70 75 80

Ser Ile Ala Asp Ala Arg Ala Gln Ala Met Arg Ala Arg Ser Glu Ala 85 90 95

Thr Glu Phe Arg Tyr Lys Tyr Gly Tyr Glu Met Pro Val Glu Ser Leu 100 105 110

Ser Arg Arg Met Ala Asn Ile Ser Gln Leu Tyr Thr Gln Arg Ala Tyr 115 120 125

Met Arg Pro Leu Gly Val Ala Leu Thr Phe Ile Gln Val Asp Phe Ala 130 135 140

Asp Glu Gly Arg Gly Pro Gln Ile Phe Lys Cys Asp Pro Ala Gly Tyr 145 150 155 160

Phe Thr Gly Val Lys Ala Val Ala Thr Gly Pro Lys Gln Gln Glu Ala 165 170 175

Thr Thr Tyr Leu Glu Lys Lys Phe Lys Lys Thr Asp Ala Val Lys Gly
180 185 190

Asp Trp Gln Lys Thr Val Glu Phe Ala Ile Ile Ala Leu Ser Ser Val 195 200 205

Ile Gly Thr Glu Phe Arg Lys Asn Asp Ile Glu Ile Gly Val Ala Thr 210 215 220

Glu Gly Glu Phe Arg Ile Leu Thr Pro Glu Glu Ile Asp Glu Arg Leu 225 230 235 240 Ile Ser Ile Ala Glu Gln Asp 245

<210> 345

<211> 968

<212> DNA

<213> Candida albicans

### <400> 345

cattgtaggt acacctgttt ttgctcaatg tacacacaca cgcaccagca gtaggaaaaa 60 aaacaaaatt aaatgaaaaa tcattttcgt tcaatattaa gcttcttaag ataaccaacc 120 aattaatatg tatgtgacat accatataaa taaagctaca aatggggata actatgtatt 180 taatgataaa tgaatggaag accagaatgt ataatgttat aagatagtga tttatattga 240 aaacaccctt aaaaaaatca accacccatc taaccgtcga attggaaatg tcaatttagt 300 tagcatcgaa aatcaacaaa gacatgggga atcatttaca tataaaataa tgagagagaa 360 ttacaaactg ctacgttatg ttttgttcat tatgtcttgt tcattatgtc ttgttcatta 420 tgtcaaaaaa tctatagtta cctacctcta catcaattta tggtctgaaa tactaacatt 480 ttttttattt atagtgaaaa atgaagattg aagttgactc cttttcaggt tctaaaatct 540 acccaggtag aggtacttta tttgtcagag gtgactctaa aatttttaga ttccaatcct 600 caaaatctgc ttctttattc caacaaagaa agaacccaag aagaatttct tggactgttt 660 tgtacagaag acaccacaaa aaaggtattt ctgaagaagc tgctaaaaaag agaaccagaa 720 agaccgtcaa gcaccaaaga gctattgtcg gtgcttcttt ggaattgatc aaagaaagaa 780 gaagtcaaaa accatctgac agaaaagctg ctagagactc taaattagct aaagacaaag 840 aagctaaaaa agctgctaaa gctgccagaa aagctgaaaa ggctaaggct gttgcttctg 900 gtgcttctgt tgtttctaaa caacaagcta aaggttcttt ccaaaaagtt aaagctacct 960 968 cccgttaa

<210> 346

<211> 155

<212> PRT

<213> Candida albicans

<400> 346

Met Lys Ile Glu Val Asp Ser Phe Ser Gly Ser Lys Ile Tyr Pro Gly

1 5 10 15

Arg Gly Thr Leu Phe Val Arg Gly Asp Ser Lys Ile Phe Arg Phe Gln 20 25 30

Ser Ser Lys Ser Ala Ser Leu Phe Gln Gln Arg Lys Asn Pro Arg Arg 35 40 45

Ile Ser Trp Thr Val Leu Tyr Arg Arg His His Lys Lys Gly Ile Ser 50 55 60

```
Glu Glu Ala Ala Lys Arg Thr Arg Lys Thr Val Lys His Gln Arg
65 70 75 80
```

Ala Ile Val Gly Ala Ser Leu Glu Leu Ile Lys Glu Arg Arg Ser Gln 85 90 95

Lys Pro Ser Asp Arg Lys Ala Ala Arg Asp Ser Lys Leu Ala Lys Asp 100 105 110

Lys Glu Ala Lys Lys Ala Ala Lys Ala Ala Arg Lys Ala Glu Lys Ala 115 120 125

Lys Ala Val Ala Ser Gly Ala Ser Val Val Ser Lys Gln Gln Ala Lys 130 135 140

Gly Ser Phe Gln Lys Val Lys Ala Thr Ser Arg 145 150 155

<210> 347 <211> 1418 <212> DNA <213> Candida albicans

<400> 347

aatctaccat taggcaatgt catataattt agattacgaa cagctgatga cgcagtttca 60 ttatctccat aatttataat ggcataccct ttggatttac ctgttaattc atcaaataat 120 aacttcacat ctaatactgg tccaactgat cttgcaatat cccgcacctg ttcttcagtg 180 taatcaaatg gaaatttacc tattgaaaca caagtgcttt tactatccat atttagtttg 240 agetgeetag tttteetgat tettggtaac aaaaaactat aattatttaa caaattateg 300 atagaatctc ttccaaagta caaccaattt cgatacattt ttttctatga gatgtttttt 360 teceetttgt etttaactaa tgtatttggg tacactacet gaacetacea gtteagttea 420 gtctaaaaaa ttttttattt attgtgtttg tttatactac cctttgcata tattactaat 480 atttatctta ccctttcatc atggatacaa aggaaataag atctaccgta tctaatcttg 540 aaaaagcagt agatgacacc accattttaa agttgttgaa tatcttgaat gatggggtta 600 aaccatccga aaaactcttg agagaaacca aagtaggtgt agctgtcaac aaattcagaa 660 gtcacgacag tgccgaaatc aacggtttag ttaaaaaaat gatcagaaat tggagagatg 720 ccgtgcaagc cgaaaagaac aacaaaaaga agttagcaat agcagctgga acaggaacag 780 gaacacette aageteagea atttegeeat cateateggg ttetggaagt acaacaceaa 840 aaccatcaga gtcaaccaca ccatcggctg cccgtaaagg tccaagaaat ccaaaaactg 900 acggtgtaaa tactcaatta tacgaaaatg ataccagaaa tgcatcggtt agtgctttat 960 acacgtcgtt agcagttgat cgtgacgatt caccaaaaca cattttgaga atagctattg 1020 aaatcgaggc tgaagtatac aagagcgaat acctgaaagt aagtgacagt tatagaaata 1080 ggttaagaag ctttaccatg aatcttcgaa ataagaaaaa cccagaatta agagaaagaa 1140 tcttatcgaa acaaatttta cctgctgcgt tcattaaaat gacccctaat gaaatggctc 1200 ctgaggcatt gaaaaaggaa attgaaaaat tgcacaaaca aaacttgttt gatgctcaag 1260 gtgctactga aaagagagca gtgaccgata gattcacttg tggcaagtgt aagcacaaga 1320 aggtcagtta ttatcaaatg caaaccagat cagcggatga gcctttaact acattttgta 1380 cttgtgaaaa ttgtggcaat agatggaagt tttcataa 1418

<210> 348

<211> 305

<212> PRT

<213> Candida albicans

<400> 348

Met Asp Thr Lys Glu Ile Arg Ser Thr Val Ser Asn Leu Glu Lys Ala

5 10

Val Asp Asp Thr Thr Ile Leu Lys Leu Leu Asn Ile Leu Asn Asp Gly
20 25 30

Val Lys Pro Ser Glu Lys Leu Leu Arg Glu Thr Lys Val Gly Val Ala 35 40 45

Val Asn Lys Phe Arg Ser His Asp Ser Ala Glu Ile Asn Gly Leu Val
50 55 60

Lys Lys Met Ile Arg Asn Trp Arg Asp Ala Val Gln Ala Glu Lys Asn 65 70 75 80

Asn Lys Lys Leu Ala Ile Ala Ala Gly Thr Gly Thr Pro 85 90 95

Ser Ser Ser Ala Ile Ser Pro Ser Ser Ser Gly Ser Gly Ser Thr Thr
100 105 110

Pro Lys Pro Ser Glu Ser Thr Thr Pro Ser Ala Ala Arg Lys Gly Pro 115 120 125

Arg Asn Pro Lys Thr Asp Gly Val Asn Thr Gln Leu Tyr Glu Asn Asp 130 135 140

Thr Arg Asn Ala Ser Val Ser Ala Leu Tyr Thr Ser Leu Ala Val Asp 145 150 155 160

Arg Asp Asp Ser Pro Lys His Ile Leu Arg Ile Ala Ile Glu Ile Glu 165 170 175

Ala Glu Val Tyr Lys Ser Glu Tyr Ser Lys Val Ser Asp Ser Tyr Arg 180 185 190

Asn Arg Leu Arg Ser Phe Thr Met Asn Leu Arg Asn Lys Lys Asn Pro 195 200 205

Lys Lys Val Ser Tyr Tyr Gln Met Gln Thr Arg Ser Ala Asp Glu Pro 275 280 285

<del>----265---</del>

Leu Thr Thr Phe Cys Thr Cys Glu Asn Cys Gly Asn Arg Trp Lys Phe 290 295 300

Ser

<210> 349 <211> 1301 <212> DNA <213> Candida albicans

<del>-260--</del>

## <400> 349

acatttattg ttggctatac tttcccagtt ttaaaggaat ggttaggcgg gtcagtctac 60 tttattttta ctgtaatgtg ttgtgccagt gtatggttta tcaagactaa agttcctgag 120 acaaagggta agcattcgta ccaggaagtc tggaattaaa cggtagttac ttccagtata 180 tatggtgttc attgtgtttt gtgtctaaag ttgcgttagg gctaaagccc taatcaatag 240 tagtgtactt tgtttgaaaa aaataataca tgcaaaatac tgcatattag aattataagg 300 gaatgaaaat gaaaaaaaa agaataattt tgtagatcgc atagtgtgag cgcgcacaca 360 cacacaaact ttgtagtgct acagtttctc tctttcccat acactctcgc agtcgcacgc 420 ttgacaaaag ttaattagaa atagaaaaat ttctcattcc ttttgagttt tccaccataa 480 tcaactagta ataaccaaca atgcctacta gattaactaa aaccagaaaa cacagaggta 540 atgtttctgg tatgtacaat ttcactaaag ttttaacgga tgaaagagta ttgaatgttc 600 aaggacatat ggagatttag agaaaacatg gaaaactaat cgaaaatgaa taatgaatgg 660 cattgatcat acaaccaaag caatgaatat gagtagtttg ggaaaccaca acatacgagt 780 tatttttcag aacaatcaag tttatcattt acagaagaca gttccatcaa tatgtccaaa 840 atactctttg aagtcaatta aagtcaaatg aacgaattac taacttattt ttttctttag 900 ccggtaaggg tagaattggt aaacacagaa agcacccggg tggtagaggt aaagctggtg 960 gtcaacatca tcacagaacc aacttggata aataccatcc aggttacttc ggtaaagttg 1020 gtatgagata cttccacaaa caacaaaacc acttctggag accagaaatc aacttggaca 1080 aattgtggac tttggttgat tctgaaaaga aagacgaata cttgagcaaa tcatctgctt 1140 ctgctgctcc agtcatcgac accttggctc acggttacgg taaagttttg ggtaaaggta 1200 gattaccaga agttccagtc attgtcaaag ccagatttgt ttctaaatta gctgaagaaa 1260 aaatcagagc tgttggtggt gttgtcgaat tagttgctta a 1301

<210> 350

<211> 149

<212> PRT

<213> Candida albicans

<400> 350

Met Pro Thr Arg Leu Thr Lys Thr Arg Lys His Arg Gly Asn Val Ser 1 5 10 15

Ala Gly Lys Gly Arg Ile Gly Lys His Arg Lys His Pro Gly Gly Arg
20 25 30

Gly Lys Ala Gly Gly Gln His His Arg Thr Asn Leu Asp Lys Tyr 35 40 45

His Pro Gly Tyr Phe Gly Lys Val Gly Met Arg Tyr Phe His Lys Gln 50 55 60

Gln Asn His Phe Trp Arg Pro Glu Ile Asn Leu Asp Lys Leu Trp Thr
65 70 75 80

Leu Val Asp Ser Glu Lys Lys Asp Glu Tyr Leu Ser Lys Ser Ser Ala 85 90 95

Ser Ala Ala Pro Val Ile Asp Thr Leu Ala His Gly Tyr Gly Lys Val 100 105 110

Leu Gly Lys Gly Arg Leu Pro Glu Val Pro Val Ile Val Lys Ala Arg 115 120 125

Phe Val Ser Lys Leu Ala Glu Glu Lys Ile Arg Ala Val Gly Gly Val 130 135 140

Val Glu Leu Val Ala 145

<210> 351

<211> 423

<212> DNA

<213> Candida albicans

```
<400> 351
gctatcaaaa tcactgttca caatggtgac agaaaacacg ttgctgcttt aagaactgtc 60
aaatctttga ttgctaactt gatcactggt gtcactaaag gttacaaata caagatgaga 120
tttgtttatg cgcatttccc aattaacgtt aacattatta aaaaagatgg tcaagattac 180
gttgaaatta gaaatttctt gggtgaaaaa agagttagag aagttaaaat ccatgaaggt 240
gtcaccatgg aaatttcttc tactcaaaag gatgaattga ttgtttctgg taactccttg 300
gaagctgttt ctcaaaatgc tgctgatatt caacaaatct gtcgtgtcag aaacaaggat 360
atccgtaaat tcttggatgg tatttatgtt tctgaaagag gtaccattgt tgaagaaatc 420
taa
```

<210> 352

<211> 140

<212> PRT

<213> Candida albicans

<400> 352

Ala Ile Lys Ile Thr Val His Asn Gly Asp Arg Lys His Val Ala Ala 1 5 10 15

Leu Arg Thr Val Lys Ser Leu Ile Ala Asn Leu Ile Thr Gly Val Thr
20 25 30

Lys Gly Tyr Lys Tyr Lys Met Arg Phe Val Tyr Ala His Phe Pro Ile 35 40 45

Asn Val Asn Ile Ile Lys Lys Asp Gly Gln Asp Tyr Val Glu Ile Arg
50 55 60

Asn Phe Leu Gly Glu Lys Arg Val Arg Glu Val Lys Ile His Glu Gly 65 70 75 80

Val Thr Met Glu Ile Ser Ser Thr Gln Lys Asp Glu Leu Ile Val Ser 85 90 95

Gly Asn Ser Leu Glu Ala Val Ser Gln Asn Ala Ala Asp Ile Gln Gln 100 105 .110

Ile Cys Arg Val Arg Asn Lys Asp Ile Arg Lys Phe Leu Asp Gly Ile
115 120 125

Tyr Val Ser Glu Arg Gly Thr Ile Val Glu Glu Ile 130 135 140

<210> 353

<211> 1655

```
<212> DNA
 <213> Candida albicans
 <400> 353
 gatgatcgct aacaaggtac ccgatttatt catcaacaaa gaagctactt catagaattg 60
 attctcaacc ttaaatagct tgtagttttt atgattggga ctaattgtgc agcatcgaac 120
 aatattgcca gttgcataga ataaatcggt attattacgg cataccaatc gattgtaggg 180
 ggttgaagct ggtgctaatt ctggggttgt tgcatttcca gtccataaac gttgaaacac 240
aaattgctgt gtaatttggt gaatgaattg atcagatgat cctatgggtg gcattgttaa 300
ggttgttcgc tacgatgtat ttctaggata atgaaggtaa gtacgaactt atgagttttg 360
atttccttcc tgatgttttt ttttccttgt tttcttttgt cgttgtagca agaagaaaaa 420
aaaaaagtgt Cttcacaagt cttggactca attttcaccc ctccacaaac tcaatttcaa 480
ttaactatca ataatccaat atgggtaaac agtatatatc taccgtcagt gcatctcagg 540
ctcataagct ggatattctt ggtgtagcta ttaccaataa gttcactgta tccgtgtcta 600
gtgatggata tgcgaaattt tgggacaaca agcaagacga agttcatctg cctaaagaat 660
ttgtccaact ggtatttata gataaaagcg gaatccatgc ggtggctgct tacgaaaatg 720
ttttgccaag ctctacattg aaagtgacat tattagcatt tgcatgtttc aatggatcta 780
tcatcttcag atattatatc aatgatgact tttcaactat cgaaagtcta actgatgata 840
taaaatcatt tgaaagcaat tgttggaccc ctggctttta tcgcgatcca gaatccaaac 900
aagactattt tattacaacc aagaccaatg gcactacaga ggttcattta ttgaatattg 960
ttgatgaaaa tgagaaggct gtaatcacat ttgaaaagtt tgggcaatta aaaggaaact 1020
cttcttcttt cccaaattct ttggctatat gtccaacaga gaataaaaaa tgtgctgtgg 1080
ggtacatcaa tggtgatgtc ttgttatatg actttgttag cttgaaattg atatacacat 1140
ttcgttcgag tgatttggtg accagtagaa attcccaatc gacgtctata cctagggtgt 1200
tggcattttc ccctggtgga accttgttgg ctgtggcaag agacaatcaa gctgctgggt 1260
caattacatt atacgacgtt gagcatggtg agaatgtggg gtctttggcc acaccctcac 1320
actcggccaa atctgttgtt ggtgggtttg cacatcaagg ctggattttg gggttgagtt 1380
ttgatgagga aggtaagcac ttggctagtt gtggatttga caaatgcata agagtctgga 1440
atttagaaac aagcgaaagg gaagcaacaa ttagtatatc tatatcagac ttagatgata 1500
ctacacataa tgatcaagac gagagtgtcg cttctggtgt tgcttttatt aaaaaggggg 1560
ttagaggtgg ctctggtggt gacagcaatg aaggattatg tgtcgtgagt ttcgatagag 1620
gaataagatg gtaccgagag gcaggaggaa tatag
                                                                   1655
<210> 354
<211> 384
<212> PRT
<213> Candida albicans
Met Gly Lys Gln Tyr Ile Ser Thr Val Ser Ala Ser Gln Ala His Lys
  1
                                     10
                                                         15
```

Ser Ser Asp Gly Tyr Ala Lys Phe Trp Asp Asn Lys Gln Asp Glu Val

Ser Asp Ile Leu Gly Val Ala Ile Thr Asn Lys Phe Thr Val Ser Val

25

30

20

40

45

His Ser Pro Lys Glu Phe Val Gln Ser Val Phe Ile Asp Lys Ser Gly 50 55 60

Ile His Ala Val Ala Ala Tyr Glu Asn Val Leu Pro Ser Ser Thr Leu 65 70 75 80

Lys Val Thr Leu Leu Ala Phe Ala Cys Phe Asn Gly Ser Ile Ile Phe 85 90 95

Arg Tyr Tyr Ile Asn Asp Asp Phe Ser Thr Ile Glu Ser Leu Thr Asp 100 105 110

Asp Ile Lys Ser Phe Glu Ser Asn Cys Trp Thr Pro Gly Phe Tyr Arg 115 120 125

Asp Pro Glu Ser Lys Gln Asp Tyr Phe Ile Thr Thr Lys Thr Asn Gly 130 135 140

Thr Thr Glu Val His Leu Leu Asn Ile Val Asp Glu Asn Glu Lys Ala
145 150 155 160

Val Ile Thr Phe Glu Lys Phe Gly Gln Leu Lys Gly Asn Ser Ser Ser 165 170 175

Phe Pro Asn Ser Leu Ala Ile Cys Pro Thr Glu Asn Lys Lys Cys Ala 180 185 190

Val Gly Tyr Ile Asn Gly Asp Val Leu Leu Tyr Asp Phe Val Ser Leu
195 200 205

Lys Leu Ile Tyr Thr Phe Arg Ser Ser Asp Leu Val Thr Ser Arg Asn 210 215 220

Ser Gln Ser Thr Ser Ile Pro Arg Val Leu Ala Phe Ser Pro Gly Gly
225 230 235 240

Thr Leu Leu Ala Val Ala Arg Asp Asn Gln Ala Ala Gly Ser Ile Thr 245 250 255

Leu Tyr Asp Val Glu His Gly Glu Asn Val Gly Ser Leu Ala Thr Pro 260 265 270

Ser His Ser Ala Lys Ser Val Val Gly Gly Phe Ala His Gln Gly Trp 275 280 285

Ile Leu Gly Leu Ser Phe Asp Glu Glu Gly Lys His Leu Ala Ser Cys

SEQL

290

295

300

Gly Phe Asp Lys Cys Ile Arg Val Trp Asn Leu Glu Thr Ser Glu Arg 305 310 315 320

Glu Ala Thr Ile Ser Ile Ser Ile Ser Asp Leu Asp Asp Thr Thr His
325 330 335

Asn Asp Gln Asp Glu Ser Val Ala Ser Gly Val Ala Phe Ile Lys Lys 340 345 350

Gly Val Arg Gly Gly Ser Gly Gly Asp Ser Asn Glu Gly Leu Cys Val 355 360 365

Val Ser Phe Asp Arg Gly Ile Arg Trp Tyr Arg Glu Ala Gly Gly Ile 370 375 380

<210> 355

<211> 1418

<212> DNA

<213> Candida albicans

<400> 355

atagagatgc ttattctggt ggttcagtta atttatatca tgtcactgaa caaggttgga 60 cttatcatgg taatttcaat gttggtgatc tcttttggga agttaaagaa aaggaacaat 120 catttgtaaa tgtagatgga taaagtaagt tgattgtata aaaaaagaat ggattcatta 180 atataaattt acaaatctga taataaacta cagtaactat tatgcaacaa ttgaaagtat 240 cctgtgtcac gtgattaggg ctcaaaagcc ctaacaggtt gcacgtgatg ttgatttatt 300 tatccctgca cacattgagt ttttttcta tggtgaaaat tattagtgag acgatgttgt 360 tegeatttta ecacacacte teacactgag tgtagteata etaacaatet teteacacta 420 tacacaaaaa aaatctttcg tttgaaaagt tttgaaaggt tcgttttctc aatagtatat 480 ccatacaata actgccaaag atggtatgtt caagtgttta aagagatcga aatgttttgt 540 tttaattgga agtgatatca taatgagaaa atatgggaaa tagagatagg accgaattaa 600 ttgaaagagt tttcttttgg ggaaatgggt tcaataagat ttttcaagat tggaatcaaa 660 ttatgtcaga tgtgaagaag agtaaatata acaagaagtt caaatcatta aaaattttat 720 gtaaatacaa cgattatcaa ttacagtcaa cgaatagaat aaaacaaaag ccagcatata 780 acagatacca gtgattcatc ccagtgatat aaataatgat attcaattaa aaacaatatt 840 aagacatttt ttatttatgt atcaacaaaa tactaactta tctttctttt aattagtctg 900 acaaatccca aaatgttatg cgtgaattac gtattgaaaa attagtttta aacatttgtg 960 ttggtgaatc cggtgataga ttaaccagag ccgccaaagt tttagaacaa ttatctggtc 1020 aaaccccagt tcaatctaaa gctagataca ctgtcagaac tttcggtatt agaagaaatg 1080 aaaaaattgc cgtccacgtt actgtcagag gtccaaaagc tgaagaaatc ttggaaagag 1140 gtttgaaagt taaagaatat caattaagat ctaaaaactt ctctgctacc ggtaactttg 1200 gtttcggtat tgatgaacat attgatttag gtatcaaata tgatccatct attggtattt 1260 acggtatgga tttctacgtt gttatgggta gagccggtgc tagagtcacc agaagaaaga 1320 gagctagatc cactattggt aactctcaca aaaccaacaa agaagatacc atccaatggt 1380 tcaagaccag atacgatgct gaagttttgg ataaataa 1418

<210> 356

<211> 174

<212> PRT

<213> Candida albicans

<400> 356 —

Met Ser Asp Lys Ser Gln Asn Val Met Arg Glu Leu Arg Ile Glu Lys

1 5 10 15

Leu Val Leu Asn Ile Cys Val Gly Glu Ser Gly Asp Arg Leu Thr Arg
20 25 30

Ala Ala Lys Val Leu Glu Gln Leu Ser Gly Gln Thr Pro Val Gln Ser 35 40 45

Lys Ala Arg Tyr Thr Val Arg Thr Phe Gly Ile Arg Arg Asn Glu Lys 50 55 60

Ile Ala Val His Val Thr Val Arg Gly Pro Lys Ala Glu Glu Ile Leu 65 70 75 80

Glu Arg Gly Leu Lys Val Lys Glu Tyr Gln Leu Arg Ser Lys Asn Phe 85 90 95

Ser Ala Thr Gly Asn Phe Gly Phe Gly Ile Asp Glu His Ile Asp Leu 100 105 110

Gly Ile Lys Tyr Asp Pro Ser Ile Gly Ile Tyr Gly Met Asp Phe Tyr 115 120 125

Val Val Met Gly Arg Ala Gly Ala Arg Val Thr Arg Arg Lys Arg Ala 130 135 140

Arg Ser Thr Ile Gly Asn Ser His Lys Thr Asn Lys Glu Asp Thr Ile 145 150 155 160

Gln Trp Phe Lys Thr Arg Tyr Asp Ala Glu Val Leu Asp Lys 165 170

<210> 357

95

```
<211> 919
<212> DNA
<213> Candida albicans
<400> 357
tattgcgtaa tcgttatcaa tgtttggatt gttgcaacgc tgggttgtgt gtatatgcga 60
gaaatgagta attgtttaaa aataatcgat gcagcagtct gccactaaat aaataaaatg 120
tcatattgac atgataagga tttatcaata ctgttaggtc tatagcccta actttaatca 180
tttacacgtg atacaaaaaa gttgtttgat cccgcacgac tatgagtacg cactcactaa 240
ttatagcctg aaaaaaaaaa tttccacata gtaaggggat tttgtatggt ggtgcgctcg 300
cctaagacgt ctgcatacat tttctaaagt cacactgata tagggatgtt gtggtagtga 360
ttgtgtgttc ccaccaaata actttgcgga cactctcata tactcaattt ttttcttaca 420
aaaatttttt ttcttctact tttttcaaga attcttcttt tacaattcaa caacatcaat 480
catgggtaaa ggtaaaccaa gagggcttaa ctctgctaga aaattaagag ttcacagaag 540
aaacaacaga tgggctgatc aagcttataa agctagatta ttaggtaccg ctttcaaatc 600
ttctccattt ggtggttcat ctcacgccaa aggtatcgtt ttggaaaaaa ttggtattga 660
atctaaacaa ccaaactctg ctatcagaaa atgtgtcaga gtccaattaa tcaaaaacgg 720
taagaaagtc actgctttcg ttccaaacga tggttgtttg aactttgttg acgaaaatga 780
cgaagtettg ttggetggtt teggtagaag aggtaaaget aagggggata tteeaggggt 840
tagattcaag gttgttaaag tttccggtgt ctctttatta gctttatgga aagaaagaa 900
agaaaagcca agatcatag
<210> 358
<211> 145
<212> PRT
<213> Candida albicans
<400> 358
Met Gly Lys Gly Lys Pro Arg Gly Leu Asn Ser Ala Arg Lys Leu Arg
  1
Val His Arg Arg Asn Asn Arg Trp Ala Asp Gln Ala Tyr Lys Ala Arg
             20
                                 25
Leu Leu Gly Thr Ala Phe Lys Ser Ser Pro Phe Gly Gly Ser Ser His
                             40
         35
Ala Lys Gly Ile Val Leu Glu Lys Ile Gly Ile Glu Ser Lys Gln Pro
                         55
Asn Ser Ala Ile Arg Lys Cys Val Arg Val Gln Leu Ile Lys Asn Gly
                     70
                                         75
Lys Lys Val Thr Ala Phe Val Pro Asn Asp Gly Cys Leu Asn Phe Val
```

Asp Glu Asn Asp Glu Val Leu Leu Ala Gly Phe Gly Arg Arg Gly Lys

90

428

105

110

Ala Lys Gly Asp Ile Pro Gly Val Arg Phe Lys Val Val Lys Val Ser 115 120 125

Gly Val Ser Leu Leu Ala Leu Trp Lys Glu Lys Lys Glu Lys Pro Arg 130 135 140

Ser

145

```
<210> 359
<211> 1164
<212> DNA
<213> Candida albicans
```

<400> 359

```
aattcatttg atctctaaga agatgaacag gccttgaatg ggaggggttg gttcgacttc 60
cacgaaatgt ggttggaaaa acttgagtcc tcaaaaaggt tctctcgtaa aaggccagag 120
ttccttactt tgcaaatgtt gaatccagtt atattcatta aagatcctat aaaatacgat 240
tattcacaat ttattatatc tttactcccg aaattcatta attgtaatcg tattgattta 300
gttatacttt gtcaaatcac cgaatcaaat caattgaatg aaattttatg tttttattat 360
caattaattc gtaatcataa gaataatgga gatactgatg gagataccga cagtttgcct 420
atgtttgatt atcgatttga aattaatata ttatttaatt tatcaacaaa aaaattgaat 480
caattatgtt taaataattg gaatcatgga tatattgcgg aaggtgataa tgataacagt 540
actaacttgt catctttgcc attgtcaata acacaaatat caaacattga aattccaaca 600
atccaatcaa gagcaaatag tagtagtgct tataatgatg aggatgataa aattactact 660
agtcggcaat atcaacaatt taaaactact gctgtaggtg gaacatttga tcatttacat 720
gatggtcata aaattttatt atcaatggca atttttttaa cttcaaataa attaattatt 780
ggtataactg gttctaattt attaattaat aaaaaattta aatctcaatt acaaactttt 840
aatcaaagac aaaatttagt tattcaattc ataaatttat tattattgag tgaaaccagt 900
gttatttttt ttgaaattta tgaaattaat gatgtttgtg gtccaactgg ttatattaat 960
gatattgata atttaataat atctcaagaa actaaatctg gtggtgaatt tgttaacaaa 1020
tttcgtaaag atcatggatt taaattatta gatattacaa taattaaagt gattggtggg 1080
aatattgaag aaaattcatg gaaaggtaaa ttaagttcaa ctgatattag agaacaagaa 1140
tataatcgat tattaaatca ataa
                                                               1164
```

```
<210> 360
```

<400> 360

Met Leu Asn Pro Val Ile Phe Ile Lys Asp Pro Ile Lys Tyr Asp Tyr

<sup>&</sup>lt;211> 322

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Candida albicans

							Ŀ				)				
1				5					10					15	
Ser	Gln	Phe	Ile 20	Ile	Ser	Leu	Leu	Pro 25	Lys	Phe	Ile	Asn	Cys 30	Asn	Arg
Ile	Asp	Leu 35	Val	Ile	Leu	Cys	Gln 40	Ile	Thr	Glu	Ser	Asn 45	Gln	Leu	Asn
Glu	Ile 50	Leu	Cys	Phe	Tyr	Tyr 55		Leu	Ile	Arg	Asn 60	His	Lys	Asn	Asn
Gly 65	Asp	Thr	Asp	Gly	Asp 70	Thr	Asp	Ser	Leu	Pro 75	Met	Phe	Asp	Tyr	Arg 80
Phe	G1u	Ile	Asn	Ile 85	Leu	Phe	Asn	Leu	Ser 90	Thr	Lys	Lys	Leu	Asn 95	Gln
Leu	Cys	Leu	Asn 100	Asn	Trp	Asn	His	Gly 105	Tyr	Ile	Ala	Glu	Gly 110	Asp	Asn
Asp	Asn	Ser 115	Thr	Asn	Leu	Ser	Ser 120	Leu	Pro	Leu	Ser	Ile 125	Thr	Gln	Ile
	Asn 130	Ile	Glu	Ile	Pro	Thr 135	Ile	Gln	Ser	Arg	Ala 140	Asn	Ser	Ser	Ser
Ala 145	Tyr	Asn	Asp	Glu	Asp 150	Asp	Lys	Ile	Thr	Thr 155	Ser	Arg	Gln	Tyr	Gln 160
Gln	Phe	Lys	Thr	Thr 165	Ala	Val	Gly	Gly	Thr 170	Phe	Asp	His	Leu	His 175	Asp

Gly His Lys Ile Leu Leu Ser Met Ala Ile Phe Leu Thr Ser Asn Lys
180 185 190

Leu Ile Ile Gly Ile Thr Gly Ser Asn Leu Leu Ile Asn Lys Lys Phe 195 200 205

Lys Ser Gln Leu Gln Thr Phe Asn Gln Arg Gln Asn Leu Val Ile Gln 210 215 220

Phe Ile Asn Leu Leu Leu Ser Glu Thr Ser Val Ile Phe Phe Glu 225 230 235 240

Ile Tyr Glu Ile Asn Asp Val Cys Gly Pro Thr Gly Tyr Ile Asn Asp 245 250 255

Ile Asp Asn Leu Ile Ile Ser Gln Glu Thr Lys Ser Gly Glu Phe

02-07-1999 [EP99870141.1] [SEQL

260 265

Val Asn Lys Phe Arg Lys Asp His Gly Phe Lys Leu Leu Asp Ile Thr 275 280 285

Ile Ile Lys Val Ile Gly Gly Asn Ile Glu Glu Asn Ser Trp Lys Gly 290 295 300

Lys Leu Ser Ser Thr Asp Ile Arg Glu Gln Glu Tyr Asn Arg Leu Leu 305 310 315 320

Asn Gln

<210> 361

<211> 1427

<212> DNA

<213> Candida albicans

<400> 361

caagagtaaa atctaaatcg ttccaatgat tttggacagc tcataggtta agaggtctac 60 atgtgttggt ggcagtagta ttggtattaa gtgggtgggt catggtatta gagagtgggt 120 gttataaaag aaggttgtga ttttttttt ccactggtgg tggtggtgct gattgtactg 180 ctgttcactt tattgtttcg tttcgttttt cctttgtttt gttgacgttg acattttttt 240 tgctgttgtt gttgtatggg aatttttgta ttagttgttc ttttttgttt tttttactt 300 gttcaaagtt tcatataata ataatattct tgttttcata ttttttttt cagaacaaga 360 acaaatacaa cttatataat ttgatcttac tcttatcccc aggtttttct atttgttttt 420 gggtttgcca ttcatatata tatatctatt attcaaatca atttgaggag tatcattaat 480 ttaaattata tcagttaaca atgtcgtatc gtggtcctaa tcaatttggt aatcaacctc 540 cacatcatgg aataccttct caacctcaac cacatattgg tccaatatct tccagcaaaa 600 gtcctttaga acaatttgaa gatgttgcta aaaaagttga agattggatc gatgattatt 660 ttaaagtctt gaaaccatac gtcccagcaa ttggtagagc atttttggtg gccactttct 720 atgaggatac tttaagaatc ttcactcaat ggaatgaaca agtttattac ttgcacaact 780 atagacacta ttggcgttgg ttgaccgttt tattcttgat caataatatg gtggttatga 840 cagttgcatc cactttagta attgccagaa aaaagaataa cattgctact attgcattga 900 tcgttgttgt tattatacaa ggtattgggt atggtttatt gtttgatgct caatttgttt 960 tgagaaactt gtccgttgtt ggagggttag tattagcatt ttccgatagt attgttagag 1020 ataaaagatc cttaaacatg ccaggtttac cgatgttgaa caatcaagac aacaaaagt 1080 atttcctttt agctggtaga attttgttag tattattatt tttgggattc gtcttttctt 1140 ctgattggtc attgggtaga gttttcatta ttataatcgg gttaacttct tgtgcttcaa 1200 ttgttgttgg ttacaagaca aagttttcag ctgctatcat gcttattgtt ttattcttat 1260 acaatgtgtt cactaaccaa ttctgggctt atgcatctca agatgctaga cgtgactttt 1320 tgagatatga attcttccaa gttttgtcaa ttgtgggagg attattgtta gtggttaatg 1380 caggtgctgg tgaattctcc atcgatgaaa agaaaaagat ttattaa 1427 <210> 362

<211> 308

<212> PRT

<213> Candida albicans

<400> 362

Met Ser Tyr Arg Gly Pro Asn Gln Phe Gly Asn Gln Pro Pro His His 1 5 10 15

Gly Ile Pro Ser Gln Pro Gln Pro His Ile Gly Pro Ile Ser Ser Ser 20 25 30

Lys Ser Pro Leu Glu Gln Phe Glu Asp Val Ala Lys Lys Val Glu Asp 35 40 45

Trp Ile Asp Asp Tyr Phe Lys Val Leu Lys Pro Tyr Val Pro Ala Ile 50 55 60

Gly Arg Ala Phe Leu Val Ala Thr Phe Tyr Glu Asp Thr Leu Arg Ile
65 70 75 80

Phe Thr Gln Trp Asn Glu Gln Val Tyr Tyr Leu His Asn Tyr Arg His
85 90 95

Tyr Trp Arg Trp Leu Thr Val Leu Phe Leu Ile Asn Asn Met Val Val
100 105 110

Met Thr Val Ala Ser Thr Leu Val Ile Ala Arg Lys Lys Asn Asn Ile 115 120 125

Ala Thr Ile Ala Leu Ile Val Val Val Ile Ile Gln Gly Ile Gly Tyr 130 135 140

Gly Leu Leu Phe Asp Ala Gln Phe Val Leu Arg Asn Leu Ser Val Val 145 150 155 160

Gly Gly Leu Val Leu Ala Phe Ser Asp Ser Ile Val Arg Asp Lys Arg 165 170 175

Ser Leu Asn Met Pro Gly Leu Pro Met Leu Asn Asn Gln Asp Asn Lys
180 185 190

Lys Tyr Phe Leu Leu Ala Gly Arg Ile Leu Leu Val Leu Leu Phe Leu
195 200 205

Gly Phe Val Phe Ser Ser Asp Trp Ser Leu Gly Arg Val Phe Ile Ile 210 215 220 02-07-1999 EP99870141.1 SEQL

Ile Ile Gly Leu Thr Ser Cys Ala Ser Ile Val Val Gly Tyr Lys Thr 225 230 235 240

Lys Phe Ser Ala Ala Ile Met Leu Ile Val Leu Phe Leu Tyr Asn Val 245 250 255

Phe Thr Asn Gln Phe Trp Ala Tyr Ala Ser Gln Asp Ala Arg Arg Asp 260 265 270

Phe Leu Arg Tyr Glu Phe Phe Gln Val Leu Ser Ile Val Gly Gly Leu 275 280 285

Leu Leu Val Val Asn Ala Gly Ala Gly Glu Phe Ser Ile Asp Glu Lys 290 295 300

Lys Lys Ile Tyr 305

<210> 363

<211> 1876

<212> DNA

<213> Candida albicans

<400> 363

aattattagt tgaaaaacaa atcataccta ttgccagagc tcaaatgaaa gtgagaatta 60 cgttatctaa aaaagcatac ttaaagactt ttcaagatga aataaaacct gttattgatc 120 aaattgtgga ggaagataac aatgggaaac aatatgagat tgttgggatt atagatccta 180 taaattatag agtcttggtc acattaattg aaaatacaga tggaagcaac aaagtcgcta 240 gacaagtgta tagaatcttt tttattaata gatgcttatc ttagttatgt tccagaacaa 360 ttcatgttag gagagagaga aacgcaatca ctcaattttt gcacaaacaa aaaacgagaa 420 gatgaagaag caaaaaagaa aattttttt tctcaaccat cttaaatcct cctacaatta 480 atttcatacc aataaagaac atgtctattg tattaccatc aggaactact gacggattta 540 aagccgtctc caaatactct gccccagtgc gtcgtccaat tgaaccagtt ggtcgttact 600 tcttagccca cgcctcaaga actttgcgtg gacacacttg gtctgaattt gaaaaattgg 660 aagctgaaaa gaatgttaaa caaatcgaag tcaacgaaga tgaggatttg ggagatgaag 720 aacaaagtga agagttatta gaacacgatc caagagaatg gaagactgcc aatttatatg 780 ctgttttagg tttatctcat ttgagaagta aagccactga agatcaaatc agaagagccc 840 acagaaaaca agttttgaaa catcatccag ataaaaaatc cgctagtggt ggattagaaa 900 acgatggatt tttcaaaatt attcaaaaag cctttgaagt tatgttggac ccagtcaaga 960 gaagacaata cgattctatt gatgttgaaa atgatccaaa accaccagct ccaaaatcca 1020 aatacgattt ctttgaagct tggggtccag tatttgaaag tgaagcaaga ttttctacta 1080 aacaaccagt tccattgttg ggaaatttag aaagcaccaa agaagaagtt gatgctttct 1140 acagtttctg gggaagattt gactcttgga agacttttga attcaaagat gaagatgttc 1200 cagatgacac tgccaacaga gatcacaaac gttatattga acgtaagaat attgcccaac 1260 agaaagaaat tgaagcaaga agatcacaag agattatcga attggtcgaa agagctcatg 1320

ctgaagatcc aagaattaaa ttgttcaaag aaaaagccaa gaaagaaaag gctgctaaga 1380 aatgggaaaa agaatctgga tcaagaaaag ctgctgaaga ggctgctgcc aaaaaggctg 1440 ctgaagaggc cgctgctaag aaagccgccg aagaagccgc cgctttgaaa gccaactcta 1500 aaaaagctaa agaagctgct aaagctgcta aaaagaaaaa caagagaaac atcagagctg 1560 ctgttaaaga caataattac tttggtgact cagctaaatc tgccgacatt gatgctgatg 1620 tcgatttgtt aatcgaaaaa ttcgacgatg tcaaattagg tgaagttgct gataaagtta 1680 aagatgctga tgctgcttca gtgaaatcta cttttgttga agttgcaaag gaattggttg 1740 gagctggttc tttagatgct tcctatttga aatattttaa ctagatttt tttttgtagg 1800

aaaataatat atatagaatg taaactatca aaaacaatta tacagaagct gaactcaaac 1860

```
<210> 364
```

<211> 427

<212> PRT

<213> Candida albicans

aagactaaac agcaag

<400> 364

Met Ser Ile Val Leu Pro Ser Gly Thr Thr Asp Gly Phe Lys Ala Val 1 5 10 15

Ser Lys Tyr Ser Ala Pro Val Arg Arg Pro Ile Glu Pro Val Gly Arg
20 25 30

Tyr Phe Leu Ala His Ala Ser Arg Thr Leu Arg Gly His Thr Trp Ser 35 40 45

Glu Phe Glu Lys Leu Glu Ala Glu Lys Asn Val Lys Gln Ile Glu Val 50 55 60

Asn Glu Asp Glu Asp Leu Gly Asp Glu Glu Gln Ser Glu Glu Leu Leu 65 70 75 80

Glu His Asp Pro Arg Glu Trp Lys Thr Ala Asn Leu Tyr Ala Val Leu 85 90 95

Gly Leu Ser His Leu Arg Ser Lys Ala Thr Glu Asp Gln Ile Arg Arg 100 105 110

Ala His Arg Lys Gln Val Leu Lys His His Pro Asp Lys Lys Ser Ala 115 120 125

Ser Gly Gly Leu Glu Asn Asp Gly Phe Phe Lys Ile Ile Gln Lys Ala 130 135 140

Phe Glu Val Met Leu Asp Pro Val Lys Arg Arg Gln Tyr Asp Ser Ile 145 150 155 160 Asp Val Glu Asn Asp Pro Lys Pro Pro Ala Pro Lys Ser Lys Tyr Asp 165 170 175

Phe Phe Glu Ala Trp Gly Pro Val Phe Glu Ser Glu Ala Arg Phe Ser 180 185 190

Thr Lys Gln Pro Val Pro Leu Leu Gly Asn Leu Glu Ser Thr Lys Glu
195 200 205

Glu Val Asp Ala Phe Tyr Ser Phe Trp Gly Arg Phe Asp Ser Trp Lys 210 215 220

Thr Phe Glu Phe Lys Asp Glu Asp Val Pro Asp Asp Thr Ala Asn Arg 225 230 235 240

Asp His Lys Arg Tyr Ile Glu Arg Lys Asn Ile Ala Gln Gln Lys Glu 245 250 255

Ile Glu Ala Arg Arg Ser Gln Glu Ile Ile Glu Leu Val Glu Arg Ala 260 265 270

His Ala Glu Asp Pro Arg Ile Lys Leu Phe Lys Glu Lys Ala Lys Lys 275 280 285

Glu Lys Ala Ala Lys Lys Trp Glu Lys Glu Ser Gly Ser Arg Lys Ala 290 295 300

Ala Glu Glu Ala Ala Ala Lys Lys Ala Ala Glu Glu Ala Ala Ala Lys 305 310 315 320

Lys Ala Ala Glu Glu Ala Ala Ala Leu Lys Ala Asn Ser Lys Lys Ala 325 330 335

Lys Glu Ala Ala Lys Ala Ala Lys Lys Lys Asn Lys Arg Asn Ile Arg 340 345 350

Ala Ala Val Lys Asp Asn Asn Tyr Phe Gly Asp Ser Ala Lys Ser Ala 355 360 365

Asp Ile Asp Ala Asp Val Asp Leu Leu Ile Glu Lys Phe Asp Asp Val 370 375 380

Lys Leu Gly Glu Val Ala Asp Lys Val Lys Asp Ala Asp Ala Ala Ser 385 390 395 400

Val Lys Ser Thr Phe Val Glu Val Ala Lys Glu Leu Val Gly Ala Gly
405 410 415

Ser Leu Asp Ala Ser Tyr Leu Lys Tyr Phe Asn 420 425

<210> 365

<211> 1178

<212> DNA

<213> Candida albicans

# <400> 365

tttcatttgt aggcattgtg tagaatgtgg actgtaggaa ggtgcaaaat atatatccaa 60 taagtcacat ctcaaattgg tagtagaaga attcaacctt gggacagtat tctctgctta 120 aaatgagttt actgatcaaa taaaatattg gctcaaccat tgaaaagcta ttcttgacac 180 tttttgcagt tttagttttg gttgtttcac aattgaaaaa aaaaatgttg agccctaata 240 gccctaattg tttgcacgtg atgtattgct ttgagtgaaa agtgtacatt tgtacacaca 300 caaacaaaaa aaaactaact tcttccccca atatgctgcg tatagtgaga aatcttcttc 360 tctcttgcca tactatcgca tttttttttt tggtaactca cgaaacctaa ttggaaagaa 420 gaaaagagaa aaaaaaaaaa ttttcagttt tcaatagatt tcagtttttg aattatacat 480 attagagaaa caagttaaca atggtatgtt cattagattt accatgagtt atggagggat 540 attcctggat gatcagaata tcataatata ggagtatact attttactgg aatcaagata 600 taattgaatt gaaataaaaa ttgaataaga ggaagcatag aatacggaat gatgattaac 660 aaatcagaaa acattgtact accgaataat gaattaggat caacatatta atatcaacta 720 ctactttaaa agaacaggaa aatgagaaga gaatcagcat gaaagttacc acctcagaaa 780 acgtcaaaga aatatacttt tggtgtggat ccatgtgtta ttactgaaat catcgaaacg 840 aatataaccc tccttttccc acattgacat aactcagaca caatctttca aataaaacat 900 gtaatactaa ctattatttt tttttcgtat aggttttagt tcaagattta ttacatccat 960 ctccagccac tgaagctaaa caacacaaat tgaaaacttt agttcaacaa ccaagatctt 1020 tetttatgga egttaaatgt caaggatgte ttaatateae caetgtttte agteaegete 1080 aaactgctgt cacttgtgac tcttgttcta ctgttttgtg taccccaact ggtggtaaag 1140 ctaaattgac tgaaggttgt tcattcagaa gaaagtaa 1178

<210> 366

<211> 82

<212> PRT

<213> Candida albicans

<400> 366

Met Val Leu Val Gln Asp Leu Leu His Pro Ser Pro Ala Thr Glu Ala
1 5 10 15

Lys Gln His Lys Leu Lys Thr Leu Val Gln Gln Pro Arg Ser Phe Phe 20 25 30

Met Asp Val Lys Cys Gln Gly Cys Leu Asn Ile Thr Thr Val Phe Ser 35 40 45

```
His Ala Gln Thr Ara Val Thr Cys Asp Ser Cys Ser Thr Val Leu Cys
50 55 60
```

Thr Pro Thr Gly Gly Lys Ala Lys Leu Thr Glu Gly Cys Ser Phe Arg
65 70 75 80

Arg Lys

<<del>210> 367</del>

```
<211> 1179
<212> DNA
<213> Candida albicans
<400> 367
tttatattgt tttttgtttc tatgtatagt aattatttaa tttgatgttc cattattcat 60
gtaattgtga tgtattttt ttgttggtcc ttctttgtgt tttgatattt tgcagctggt 120
cggatcatac aaatttggaa ggagctacca aagcagaaat gatagcaaaa aaaaacagga 180
gacttttgga atttcttggt ttagggctct agccctaatg cctgtatata attgattgat 240
gctttaaaat ttgtgcggtg aaaaacagaa gcgaaaaaaa aagagctgga aagtgtgatt 300
aacaacaacg agggaagaaa aaaaacttaa ttaggtgaac aaaaattaat gtgtgagcgt 360
gtacatgcac aattttcaat tggttattct cacgcactaa caaactcaac ctccattgtg 420
agaaaagaga tttttctttg aaaaaaaatt tacatacata tcttagttca ctagtaactt 480
tcgaagacaa catagacaag atgggtatgt tccagtatag caatattcaa caatgagata 540
caagatattc ctccaataca aagagattaa tggtgttttt gatagagtgt aaagccatgt 600
gtttaataag aatgatttgt caagggataa gatgaaaagt ggaaattgtc caaacaacat 660
tgaaacgaat tttctgggat agcgttcaaa caatcaaacc tatccaggtt tcttatgaca 720
attaagaaaa caagtcaata ttatcaaaga tggagtttta taaggagaga tattgaaaca 780
tcaaattgtt gaaagaaata gccaactatt tggcaccatt tcctgtgatc atagaaatac 840
taacaatcct cttgtgttat agttaacgtt ccaaaaacta gaaagaccta ctgtaaaggt 900
aaagaatgcc gtaaacatac ccaacacaaa gttacccaat acaaagctgg taaagcttct 960
ttgttcgctc aaggtaaaag aagatatgac agaaaacaat ccggttatgg tggtcaaacc 1020
aaacaaattt tccacaagaa agccaagact accaaaaaag ttgttttgag attggaatgt 1080
gttgtctgta aaaccaaggc tcaattacca ttgaaaagat gtaaacattt cgaattgggt 1140
ggtgacaaaa aacaaaaagg tcaagcttta caattttaa
                                                                  1179
```

```
<210> 368
```

<211> 106

<212> PRT

<213> Candida albicans

## <400> 368

Met Val Asn Val Pro Lys Thr Arg Lys Thr Tyr Cys Lys Gly Lys Glu
1 5 10 15

```
Cys Arg Lys His Thr Gln His Lys Val Thr Gln Tyr Lys Ala Gly Lys
20 25 30
```

Ala Ser Leu Phe Ala Gln Gly Lys Arg Arg Tyr Asp Arg Lys Gln Ser 35 40 45

Gly Tyr Gly Gln Gln Thr Lys Gln Ile Phe His Lys Lys Ala Lys Thr
50 55 60

Thr Lys Lys Val Val Leu Arg Leu Glu Cys Val Val Cys Lys Thr Lys
65 70 75 80

Ala Gln Leu Pro Leu Lys Arg Cys Lys His Phe Glu Leu Gly Gly Asp 85 90 95

Lys Lys Gln Lys Gly Gln Ala Leu Gln Phe 100 105

<210> 369

<211> 3583

<212> DNA

<213> Candida albicans

<400> 369

gtgcgaataa tacaagcgat cgtttgaata atccaatgat tagctctgat caacgaattc 60 aaaattttcc cctcaaggac gtagtacaac aagagatgag aaggggtttg aactgtttga 120 tttcagagga gatgacggtg caaaaagatc taccaccaaa gaagcacata attttttgaa 180 tgattcacga aagagttcca ttagtgatat ctcaagccaa aaattttcaa gggaaagttc 240 ccgtaatgtt tcacgggaaa gcagcaggcg aagttcaata ataaaaattg atcatcatac 300 taatgtggac gtgtcaacga aaccggaaaa tataaatagc agagataaca aaactgagaa 360 gaatatgact ttgagttcag agtcaaccaa accgagtgtc gaagaagtta gtaaatcttt 420 gaaaccaaca attaccaaga aaacttcatt cactgattac ttaaaatctg ccaaaactaa 480 ggctaaagaa gaaaaggtaa caatcgaaaa aagtgacaaa acgattaatt cagaagaacg 540 aaaaacggaa ccaattcaac aaagcgagca acttttaacg gacaaaaagg ataataaact 600 ggaacccaat agtgaagtaa atttgaaaga caataatgac gacctgaaag ctacagctgg 660 ctgtgcccta ggacctgata agaatactgg aaaaaacgat tcagataaat cagaaacgac 720 tcaaccaaaa cttgcccgct cagaatcatt tgccgataca tcattattgt ctccagttaa 780 tgaaagtgat actgatttca atttcaatga gttagcagag ataccggaag caaaggatgg 840 ttcagttgtg gcagcaaatg tgctggagaa cattgatgag aatgaaaata tttcagaagc 900 tgaaactgtt atagcagatg accttccacg tctcgatgaa ggaaagaaac ttttacgtga 960 acaaacagca gatgtaaaga gacataaatt gaagaaaaca aaattgaata ctatttttag 1020 ttccgatgaa gaggaggagg agattcaaga accagatttc aaactccaag aaccagaaaa 1080 actaccagaa gatgaccaac atcctgattt tcaaaattca aaagcaacaa cagaaatcag 1140 taacgataaa acagaagtaa ataagccaga agtgaaagag gttggcgaga aagagagaaa 1200 tcaccaacta gaagatagat taccaattaa aaaagagaaa atgcggtcgg agaatgcaaa 1260 gacatctgaa aacggtgtca gttcaaaatc agaatctaag atttcaaagt cgaagaaact 1320

```
accttacaaa gttaaacgtg attcaagtgg tcgatcatta ttacaacgag cttgcaagaa 1380
gggtaatttt gcagatgtcc aagattacat agagaggggt gctagtgcca atgaaaagga 1440
tttctgtgga ttcacatgct tgcatgaggc agcattggaa ggtcatacac aaattgtgaa 1500
atatctcatc gaaaatggtg ctaatgtaaa tgcgaaagct gatgaagctg gtgattccga 1560
gacaccattg atcgatgcag ctgaaaacaa acatcttgat tgtgttaaag tgttactaga 1620
aaatgatgct gatcctacta ttttcaatat tgatggattc accgctttaa ctaagattta 1680
taatgaacat gaaggagaag aaggatatga tgaaattatt caagttctag aagaagcaac 1740
tgcaaactat aacagtcgtt taccaagaga agttcagttt gtatcagatg ctcctattgg 1800
tagtggacca atcatggagg atccaaatga caactatttt gcagagctaa ttaaaggtaa 1860
gggaatatat aaatacgctg cagagaattc aaaagagaaa acagcagaat attttgttgc 1920
tggtcataat ttagaaggaa aaccagatat tttaatttta gctgctagaa acggccacac 1980
agaactigit gatattatac tagggetaaa eeetacaeet titaatatig atacegagte 2040
aagttgtggt gttactgcat tattggccag tattggacgc ggtcattttg aagttgtcga 2100
tttagaaatt gcacaacact cgccacattt tgattcacgg gaagttagtg taataatgaa 2220
attcatggag aagaagagtg gaaccaaaat tttgtctggt attccgtcaa gggtagtatc 2280
tegtgcaaca tetegtgcae ettetgttee egtateatea gacgaagatg aegtagtgga 2340
agaaaaagaa attacagcac acacagaaaa taaaagtgct gaaaagaagc tggaggataa 2400
aatcacgaaa acagttaatg agcatgtcag caatcggaaa ccccacgagt ctacaggtcg 2460
aaaacttgaa aagactcata gcaatgagga aagaaagcgg aaacgtgaat ggtcagacga 2520
tgaacctaaa gaaccacatc ttttgaagaa gtcaaagtct gatttgaaat tgaaatcact 2580
acatagagaa ttcacttctg atgatcacca caccagtgaa agccattcag attcttttgc 2640
agaaaaaaga aaacatttat cggcaacgcc acctgctcct ccgccaccac caccaccacc 2700
accttctcaa gcagttatca aggcacaaga agaacaaaaa atcaaagatg ctgaagaagc 2760
tagattgtgg caagaaaagg tcgaagctaa aaagagggct cgaagagaaa tgtttttaaa 2820
gtccgagaaa gagaaagagc agaaacggaa agaagaagaa gaattgcgag cacaagaaga 2880
aaaacgaatt gccaaagcaa aacaggaaga acaggagaga cttgctagag aagcagaaga 2940
gaaatcgaaa gaactagaag aaaagaaagt ggggttgcga caacagttga cattagatca 3000
ttatccggtt ggattgcgtt attgcaagtt tgatggaaac ccaaatatct cggctgttga 3060
taaatttttg cctttctatg tatttgtaat cgacgataaa aagtatgctg ttgatttgca 3120
agtctccttg atcacgtcaa cggttgtcag caaggttatc aatactgtac aacctcatca 3180
gaagagagaa ataaatgcaa ctgaaaaaag caaattgtgg aagctctttt tcaagtttat 3240
tggtattgat cctaggaatc caaattgtga tcaaagaagc tcaataacaa atggtcaaaa 3300
acagtttcaa aatttgttgc ttcattttgt ggaggtagat ttagctgaag aatttttaaa 3360
ggaatttcca gaagtacact caaaagcaaa agataaccaa attgatgtta gtttagagtc 3420
tttgagtggg ttttctgatt gcgtcaagga tgatataata gttgatggaa atcttgaaat 3480
tgatattgat tccaagaaaa tcgaaaagtt tattcctcct catttgaata ctaggaaaga 3540
                                                                 3583
cattattagg actgtcagta ctitagcaca cccactatgg tga
```

```
<210> 370
<211> 1072
<212> PRT
<213> Candida albicans
<400> 370
Met Thr Leu Ser Ser Glu Ser Thr Lys Pro Ser Val Glu Glu Val Ser
```

10

Lys	Ser	Leu	Lуs 20	Pro	Thr	116	THE	ьуs 25	ьуs	Inr	ser	Pne	30	Asp	туг
Leu	Lys	Ser 35	Ala	Lys	Thr	Lys	Ala 40	Lys	Glu	Glu	Lys	Val 45	Thr	Ile	Glu
Lys	Ser 50	Asp	Lys	Thr	Ile	Asn 55	Ser	Glu	Glu	Arg	Lys 60	Thr	Glu	Pro	Ile
Gln	Gln	Ser	Glu	Gln	Leu	Leu	Thr	Asp	Lys	Lys	Asp	Asn	Lys	Ser	Glu
65					70					75					80
Pro	Asn	Ser	Glu	Val 85	Asn	Leu	Lys	Asp	Asn 90	Asn	Asp	Asp	Ser	Lys 95	Ala
Thr	Ala	Gly	Cys 100	Ala	Leu	Gly	Pro	Asp 105	Lys	Asn	Thr	Gly	Lys 110	Asn	Asp
Ser	Asp	Lys 115	Ser	Glu	Thr	Thr	Gln 120	Pro	Lys	Leu	Ala	Arg 125	Ser	Glu	Ser
Phe	Ala 130	Asp	Thr	Ser	Leu	Leu 135	Ser	Pro	Val	Asn	Glu 140	Ser	Asp	Thr	Asp
Phe 145	Asn	Phe	Asn	Glu	Leu 150	Ala	Glu	Ile	Pro	Glu 155	Ala	Lys	Asp	Gly	Ser 160
Val	Val	Ala	Ala	Asn 165	Val	Ser	Glu	Asn	Ile 170	Asp	Glu	Asn	Glu	Asn 175	Ile
Ser	Glu	Ala	Glu 180	Thr	Val	Ile	Ala	Asp 185	Asp	Leu	Pro	Arg	Leu 190	Asp	Glu
Gly	Lys	Lys 195	Leu	Leu	Arg	Glu	Gln 200	Thr	Ala	Asp	Val	Lys 205	Arg	His	Lys
Leu	Lys 210		Thr			Asn 215	Thr	Ile	Phe	Ser	Ser 220	Asp	Glu	Glu	Glu
Glu 225	Glu	Ile	Gln	Glu	Pro 230	Asp	Phe	Lys	Leu	Gln 235	Glu	Pro	Glu	Lys	Leu 240
Pro	Glu	Asp	Asp	Gln 245	His	Pro	Asp	Phe	Gln 250	Asn	Ser	Lys	Ala	Thr 255	Thr
Glu	Ile	Ser	Asn 260	Asp	Lys	Thr	Glu	Val 265	Asn	Lys	Pro	Glu	Val 270	Lys	Glu

Val	Gly	Glu	Lys	Glu	Arg	Asn	His	Gln	Leu	Glu	Asp	Arg	Leu	Pro	IIe
		275					280					285			

Lys Lys Glu Lys Met Arg Ser Glu Asn Ala Lys Thr Ser Glu Asn Gly 290 295 300

Val Ser Ser Lys Ser Glu Ser Lys Ile Ser Lys Ser Lys Leu Pro 305 310 315 320

Tyr Lys Val Lys Arg Asp Ser Ser Gly Arg Ser Leu Leu Gln Arg Ala

Cys Lys Lys Gly Asn Phe Ala Asp Val Gln Asp Tyr Ile Glu Arg Gly 340 345 350

Ala Ser Ala Asn Glu Lys Asp Phe Cys Gly Phe Thr Cys Leu His Glu 355 360 365

Ala Ala Leu Glu Gly His Thr Gln Ile Val Lys Tyr Leu Ile Glu Asn 370 375 380

Gly Ala Asn Val Asn Ala Lys Ala Asp Glu Ala Gly Asp Ser Glu Thr 385 390 395 400

Pro Leu Ile Asp Ala Ala Glu Asn Lys His Leu Asp Cys Val Lys Val 405 410 415

Leu Leu Glu Asn Asp Ala Asp Pro Thr Ile Phe Asn Ile Asp Gly Phe 420 425 430

Thr Ala Leu Thr Lys Ile Tyr Asn Glu His Glu Gly Glu Glu Gly Tyr
435 440 445

Asp Glu Ile Ile Gln Val Leu Glu Glu Ala Thr Ala Asn Tyr Asn Ser 450 455 460

Arg Leu Pro Arg Glu Val Gln Phe Val Ser Asp Ala Pro Ile Gly Ser
465 470 475 480

Gly Pro Ile Met Glu Asp Pro Asn Asp Asn Tyr Phe Ala Glu Leu Ile 485 490 495

Lys Gly Lys Gly Ile Tyr Lys Tyr Ala Ala Glu Asn Ser Lys Glu Lys 500 505 510

Thr Ala Glu Tyr Phe Val Ala Gly His Asn Leu Glu Gly Lys Pro Asp 515 520 525

Ile	Leu 530	Ile	Leu	Ala	Ala	Arg 535	Asn	Gly	His	Thr	Glu 540	Leu	Val	Asp	Ile
Ile 545	Leu	Gly	Leu	Asn	Pro 550	Thr	Pro	Phe	Asn	Ile 555	Asp	Thr	Glu	Ser	Ser 560
Cys	Gly	Val	Thr	Ala 565	Leu	Leu	Ala	Ser	Ile 570	Gly	Arg	Gly	His	Phe 575	Glu
Val	Val	Asp	Ser	Leu	Leu	Ser	Lys		Ala	Asp	Pro	Phe		Thr	Arg
			580					585					590		
Lys	Lys	Asp 595	Gly	Leu	Asn	Ala	Leu 600	Glu	Ile	Ala	Gln	His 605	Ser	Pro	His
Phe	Asp 610	Ser	Arg	Glu	Val	Ser 615	Val	Ile	Met	Lys	Phe 620	Met	Glu	Lys	Lys
Ser 625	Gly	Thr	Lys	Ile	Leu 630	Ser	Gly	Ile	Pro	Ser 635	Arg	Val	Val	Ser	Arg 640
Ala	Thr	Ser	Arg	Ala 645	Pro	Ser	Val	Pro	Val 650	Ser	Ser	Asp	Glu	Asp 655	Asp
Val	Val	Glu	Glu 660	Lys	Glu	Ile	Thr	Ala 665	His	Thr	Glu	Asn	Lys 670	Ser	Ala
Glu	Lys	Lys 675	Ser	Glu	Asp	Lys	Ile 680	Thr	Lys	Thr	Val	Asn 685	Glu	His	Val
Ser	Asn 690	Arg	Lys	Pro	His	Glu 695	Ser	Thr	Gly	Arg	Lys 700	Leu	Glu	Lys	Thr
His 705	Ser	Asn	Glu	Glu	Arg 710	Lys	Arg	Lys	Arg	Glu 715	Trp	Ser	Asp	Asp	Glu 720
Pro	Lys	Glu	Pro	His 725		Leu	Lys	Lys	Ser 730		Ser			Lys 735	Leu
Lys	Ser	Leu	His 740	Arg	Glu	Phe	Thr	Ser 745	Asp	Asp	His	His	Thr 750	Ser	Glu
Ser		Ser 755	Asp	Ser	Phe	Ala	Glu 760	Lys	Arg	Lys	His	Leu 765	Ser	Ala	Thr
Pro	Pro 770	Ala	Pro	Pro		Pro 775	Pro	Pro	Pro	Pro	Pro 780	Ser	Gln	Ala	Val

Ile 785	Lys	Ala	Gln		Glu 790	Gln	Lys	Ile	Lys	Asp 795	Ala	Glu	Glu	Ala	Arg 800
Leu	Trp	Gln	Glu	Lys 805	Val	Glu	Ala	Lys	Lys 810	Arg	Ala	Arg	Arg	Glu 815	Met
Phe	Leu	Lys	Ser 820	Glu	Lys	Glu	Lys	Glu 825	Gln	Lys	Arg	Lys	Glu 830	Glu	Glu
Glu	Leu	Arg	Ala	Gln	Glu	Glu		Arg	Ile	Ala	Lys	Ala 845	Lys	Gln	Glu
		835					840					043			
Glu	Gln 850	Glu	Arg	Leu	Ala	Arg 855	Glu	Ala	Glu	Glu	Lys 860	Ser	Lys	Glu	Leu
Glu 865	Glu	Lys	Lys	Val	Gly 870	Leu	Arg	Gln	Gln	Leu 875	Thr	Leu	Asp	His	Tyr 880
Pro	<b>Val</b>	Gly	Leu	Arg 885	Tyr	Cys	Lys	Phe	Asp 890	Gly	Asn	Pro	Asn	Ile 895	Ser
Ala	Val	Asp	Lys 900	Phe	Leu	Pro	Phe	Tyr 905	Val	Phe	Val	Ile	Asp 910	Asp	Lys
Lys	Tyr	Ala 915	Val	Asp	Leu	Gln	Val 920	Ser	Leu	Ile	Thr	Ser 925	Thr	Val	Val
Ser	Lys 930	Val	Ile	Asn	Thr	Val 935	Gln	Pro	His	Gln	Lys 940	Arg	Glu	Ile	Asn
Ala 945	Thr	Glu	Lys	Ser	Lys 950	Leu	Trp	Lys	Leu	Phe 955	Phe	Lys	Phe	Ile	Gly 960
Ile	Asp	Pro	Arg	Asn 965	Pro	Asn	Cys	Asp	Gln 970	Arg	Ser	Ser	Ile	Thr 975	Asn
Gly	Gln	Lys	Gln 980	Phe	Gln	Asn	Leu	Leu 985		His	Phe	Val	Glu 990	Val	Asp
Leu	Ala	Glu 995	Glu	Phe	Leu		Glu 1000	Phe	Pro	Glu	Val	His 1005		Lys	Ala
	Asp 1010		Gln	Ile		Val 1015		Leu	Glu	Ser	Leu 1020		Gly	Phe	e Ser
													_	_	_

1030

Asp Cys Val Lys Asp Asp Ile Ile Val Asp Gly Asn Leu Glu Ile Asp

1035

1040

SEQL

```
Ile Asp Ser Lys Lys Ile Glu Lys Phe Ile Pro Pro His Leu Asn Thr
1045 1050 1055
```

Arg Lys Asp Ile Ile Arg Thr Val Ser Thr Leu Ala His Pro Leu Trp 1060 1065 1070

```
<210> 371
```

<211> 659

<212> DNA

<213> Candida albicans

<400> 371

tagtttcttt tggcggtata tcctcactca tgattgatgt atttgccac taaaaagaaa 60 accaatgaaa caataagatt gattgataga tttggttgta attaatctga tgctttcaac 120 atttgtttt gttcgtgct cggtcattga ggttgggtaa attgttttc ttttgagaat 180 tgtgggcatg caatgtcgca tgcaaatatg atgtcgctca attgcgacat actacttagg 240 gctatagacc tattgcacgt gcgttagttt taaacctaaa aaaacaattt tgtgcagtcg 300 tgcaccattc gttctatttt tctactgtga ttgacgtaca aaccttcaca gttcacgac 360 acttttgtcg attctcaaac ctctctcaca ggcttggtag gaatgaaaaa aattttggta 420 aaggcgaaaa aaaaaaaaa taaacttgaa tattttggaa tccccttttt gattactaca 480 atagattaaa gtaactaaag atgattgaac catccttgaa agctttagct tcaaaaataca 540 gtcgtaagag aaagtgtggt cacaccaatc aattgagacc aaagaagaaa ttgaagtag 659

<210> 372

<211> 52

<212> PRT

<213> Candida albicans

<400> 372

Met Ile Glu Pro Ser Leu Lys Ala Leu Ala Ser Lys Tyr Asn Cys Glu

1 10 15

Lys Ser Ile Cys Arg Lys Cys Tyr Ala Arg Leu Pro Pro Arg Ala Thr 20 25 30

Asn Cys Arg Lys Arg Lys Cys Gly His Thr Asn Gln Leu Arg Pro Lys
35 40 45

Lys Lys Leu Lys 50

```
<210> 373
<211> 2297
<212> DNA
<213> Candida albicans
```

### <400> 373

```
actcagcctt ttgaaattca cttttggtaa aaggatatgt aattgaatcc gcataatgga 60
ctttctctaa ctccatattg ttgttcctac cctcttgcga tagagacttt gatcgtgaca 120
ttttgtaagt ggtagaggta tgggtaagat tattttcacc accgttttcc tttagcaaat 180
agtctaaagc attttctgaa ttgttgagtt tcggtacatc attagctatc ttttgtacaa 240
ttgatttgct tctacctcta gtcatggaaa taacaagtta acagcgtgtt ttgttggttt 300
ttttgtcttg ttctaaaaat gaaacagcgc aatgattctt ttcatagttt ttttttcca 420
tacataactt ctgacgcgtg cactatatct gctaacatac tcggcaacaa caaagaaaga 480
gaatttgaac taatccgaag atgaacgacc ccagagatga acaaattgac tccgatgatg 540
tattaacaga agattcatct gatgaattga aggacttggt gcaggagttt gaattgaaat 600
atgcagaatt aaagaagaac aaagccttga aaaaacgtcg ttcacagtca ccactggaag 660
acatgctgaa taaacagaaa ccccatcaac cggaggtccc cagaacccca gaaaaagcta 720
ctcgggattc caagataaaa gaatcaaact tcttaaacaa actctacgag acaagtaata 840
agcatgataa agaagatgcc cataaaatag atttcagtaa aagaaggttt gagtttcaat 900
tggataagta tacttttacg ccaaaagacg ttgttgatga tttagaaccg atatccaagc 960
tttacttgcg cagaagatac cttgcccaac tgcaaattgc cgacattatt gccgaaactg 1020
atagcaacat gaaattcctt aaaattgaca agtttttagc gaaaacccat aaatcaaaca 1080
actatgctga acccaagtat tgtaattggt gtcttgttgc ctttgtggtg cgcaaggacc 1140
cagtgcaggt tgctgccaac aactctaaat acatcaagtt gaaggttgga aattttatga 1200
attctgtaga cttgatgctt tttgataagg cttttcagaa aaacgggaaa atccaaccgg 1260
gtgatttatt attcattttg aatccattga tcaacaagta tgaaatacaa gttggcaagg 1320
gccagtttca gtctgggttc aatttgaaag tagaaaacac caatgtgtca agtattttag 1380
agataggatc gttaagagat tttgggtttt gcaagtttac tagaaagcta gataatagca 1440
gatgcaaaag ggccatcaac acaaggacgc aagaattctg cgatatacat ttggatatga 1500
agtttaagtc tagtacaaga atggaattaa atggaagtgt ttcgattaga tccccgcaaa 1560
aaaacaagaa aaagatgtac atgaataaaa atgggtctgg ctttattaag caatataacg 1620
aggagagtac tgttatagga acaagctacg gttctcctct tgatccaaaa aggtaccaag 1680
acccgaaagt actccagaac caaatcaagc ggcgcaaact aatagacgac aaggcaaagg 1740
agatgettga acaaaageta tegaaactag gtteggette attattgaac aatttacaat 1800
tatctaagaa agaagcaaca gataaacttg cgagtgatcg tctgaagagc aaaggattca 1860
ccaacacaat gatttcacat attgggtttg atccaacagg tacatctttg aaccaaata 1920
gtaccctgct tggcagcaag ctgatggaaa aatctcgagc acgggaattg catgatttga 1980
gtgttgaaac atctggtcat aagtcgttgt cttcgtctaa acaagatcgc cagtcgaaag 2040
ttgcaaaatg gaacacgaat atcagaactt tacaaaatta cgatcgacga gttgccagcc 2100
actccttgag tacctctcgt cggttacaga atcttgtagg aaagcaaaca cacgctacac 2160
tagtagataa aaggaaaaga gtggttgtct cagatgatga acaaccgggg atggaagaag 2220
atgaagaaga cattgaaatc cagtttgacg acgaaaagtc taagatgtcc tacatgaaaa 2280
                                                               2297
tgactggagc caggtaa
```

<210> 374

<211> 598

<212> PRT

<213> Candida albicans

<400> 374

Met Asn Asp Pro Arg Asp Glu Gln Ile Asp Ser Asp Asp Val Leu Thr

1 5 10 15

Glu Asp Ser Ser Asp Glu Leu Lys Asp Leu Val Gln Glu Phe Glu Leu 20 25 30

Lys Tyr Ala Glu Leu Lys Lys Asn Lys Ala Leu Lys Lys Arg Arg Ser 35 40 45

Gln Ser Pro Ser Glu Asp Met Ser Asn Lys Gln Lys Pro His Gln Pro 50 55 60

Glu Val Pro Arg Thr Pro Glu Lys Ala Lys Val His Leu Asp Lys Val 65 70 75 80

Val Glu Glu Pro Lys Gln Arg Ile Phe Thr Lys Lys Glu Pro Arg Asp 85 90 95

Ser Lys Ile Lys Glu Ser Asn Phe Leu Asn Lys Leu Tyr Glu Thr Ser 100 105 110

Asn Lys His Asp Lys Glu Asp Ala His Lys Ile Asp Phe Ser Lys Arg 115 120 125

Arg Phe Glu Phe Gln Leu Asp Lys Tyr Thr Phe Thr Pro Lys Asp Val 130 135 140

Val Asp Asp Leu Glu Pro Ile Ser Lys Leu Tyr Leu Arg Arg Tyr 145 150 155 160

Leu Ala Gln Ser Gln Ile Ala Asp Ile Ile Ala Glu Thr Asp Ser Asn 165 170 175

Met Lys Phe Leu Lys Ile Asp Lys Phe Leu Ala Lys Thr His Lys Ser 180 185 190

Asn Asn Tyr Ala Glu Pro Lys Tyr Cys Asn Trp Cys Leu Val Ala Phe 195 200 205

Val Val Arg Lys Asp Pro Val Gln Val Ala Ala Asn Asn Ser Lys Tyr

Ile Lys Leu Lys Val Gly Asn Phe Met Asn Ser Val Asp Leu Met Leu Phe Asp Lys Ala Phe Gln Lys Asn Gly Lys Ile Gln Pro Gly Asp Leu Leu Phe Ile Leu Asn Pro Leu Ile Asn Lys Tyr Glu Ile Gln Val Gly Lys Gly Gln Phe Gln Ser Gly Phe Asn Leu Lys Val Glu Asn Thr Asn Val Ser Ser Ile Leu Glu Ile Gly Ser Leu Arg Asp Phe Gly Phe Cys Lys Phe Thr Arg Lys Leu Asp Asn Ser Arg Cys Lys Arg Ala Ile Asn Thr Arg Thr Gln Glu Phe Cys Asp Ile His Leu Asp Met Lys Phe Lys Ser Ser Thr Arg Met Glu Leu Asn Gly Ser Val Ser Ile Arg Ser Pro Gln Lys Asn Lys Lys Met Tyr Met Asn Lys Asn Gly Ser Gly Phe Ile Lys Gln Tyr Asn Glu Glu Ser Thr Val Ile Gly Thr Ser Tyr Gly Ser Pro Leu Asp Pro Lys Arg Tyr Gln Asp Pro Lys Val Leu Gln Asn Gln Ile Lys Arg Arg Lys Leu Ile Asp Asp Lys Ala Lys Glu Met Leu Glu Gln Lys Leu Ser Lys Leu Gly Ser Ala Ser Leu Leu Asn Asn Leu Gln Leu Ser Lys Lys Glu Ala Thr Asp Lys Leu Ala Ser Asp Arg Ser Lys Ser Lys Gly Phe Thr Asn Thr Met Ile Ser His Ile Gly Phe Asp Pro Thr Gly Thr Ser Leu Asn Gln Asn Ser Thr Ser Leu Gly Ser Lys

Arg Arg Val Ala Ser His Ser Leu Ser Thr Ser Arg Arg Leu Gln Asn 530 540

Leu Val Gly Lys Gln Thr His Ala Thr Leu Val Asp Lys Arg Lys Arg 545 555 5560

Val Val Val Ser Asp Asp Glu Gln Pro Gly Met Glu Glu Asp Glu Glu 565 570 575

Asp Ile Glu Ile Gln Phe Asp Asp Glu Lys Ser Lys Met Ser Tyr Met 580 585 590

Lys Met Thr Gly Ala Arg 595

<210> 375 <211> 1499 <212> DNA <213> Candida albicans

<400> 375

tttcttcatc aacttgctca aatgtcttg aaagtgaaag cttactttt tggtttcgtt 60 taggtggcat caaggtaaag gagggagttt tggtatagtt aggtttttga cttatttcct 120 tttttgagta gatataacag aactaccaaa agtgagccca catctgttaa tcttgaaaag 180 caaaattgag aaaaccattt atgcaagtcg tgtactggtg atattcttgg tcaattgcta 240 cacttttgta atgaatactg taatgtagcc gacgtgggtt gaanaatata tatttaagta 300 tatagaatca ggtcaatata aaatgtttga aatataacaa aatgtttcaa tgtaaactga 360 tggttaaggg attataaatc aaactgagta gtgcttttgt tcctaaaaaa cccatcgtgg 420 tggtaacgtc aggagaccgc gacatcaaat ggaattcaca caatcagtct ccgaatttat 480 tacccttgaa acttaactta atgaaagagc cttccatcac tactaccttt gttgaggtta 540 cagacaaact tcctacaaag cctccgtgca gggtgtttt caagaatgag tatgagcagc 600 cctccggcag tgtcaaatta agaggcatgg gacacttggt tggccagtct atagacggc 660 ccagaaaact tggcaaatcg aacgtagcag tttttcgtc atctggtgt ttgcctgaaa 780 gttcgaagcc aactgttata gaaaagttga aatccttggg tgcagatgc attattcatg 840

```
ggaaacattg gggagaggcc gataactatt taactgattt tgttataaa aatcttgaca 900 aaacagtcta tccggtctat tgtcacctt ttgatgaccc attgttgtg gagggtcata 960 gtaagatcat cacggaaatc atcgatcaaa agcaattacc caactttgat aaagttaagg 1020 gggtcatttg ttcggtagga gggggtggct tatacaacgg aatagttgaa ggtttggaaa 1080 atcataagga gataccagtg ttggcaattg aaactaaaca agcggccacg tttcacgagg 1140 cggtcaaaga aggtaaagtt gttcatttac aaaaagtgca aactttggcc acttcttgg 1200 cttcgccgta ccttctcc aaggcattag caaactatat tgagcgtcct acagttcttg 1260 ctgaaattga tgacttggac gctgttaaag gtgttgttga tgtatacgac cattctggat 1320 atatggttga gcctgcatgt ggtgcatccg ttgcatcagt gatgcacagg caagatttat 1380 tgaatacaa caagtatat agtccagat atatatcat tgttgtcata tgtggtggat 1440 cggctatca caagtatat atagacgaat atagaagttt attagaaaaa gactcttga 1499
```

```
<210> 376
```

<211> 332

<212> PRT

<213> Candida albicans

<400> 376

Met Lys Glu Pro Ser Ile Thr Thr Phe Val Glu Val Thr Asp Lys

1 5 10 15

Leu Pro Thr Lys Pro Pro Cys Arg Val Phe Phe Lys Asn Glu Tyr Glu 20 25 30

Gln Pro Ser Gly Ser Val Lys Leu Arg Gly Met Gly His Leu Val Gly
35 40 45

Gln Ser Ile Asp Val Ala Arg Lys Leu Gly Lys Ser Asn Val Ala Val 50 55 60

Phe Ser Ser Ser Gly Gly Asn Ala Gly Leu Ala Ala Ala Tyr Ala Ser 65 70 75 80

Gln Phe Phe Gly Val Ser Cys Thr Val Val Leu Pro Glu Ser Ser Lys 85 90 95

Pro Thr Val Ile Glu Lys Leu Lys Ser Leu Gly Ala Asp Val Ile Ile
100 105 110

His Gly Lys His Trp Gly Glu Ala Asp Asn Tyr Leu Thr Asp Phe Val 115 120 125

Ile Lys Asn Leu Asp Lys Thr Val Tyr Pro Val Tyr Cys His Pro Phe 130 135 140

Ile Asp Gln Lys Gln Leu Pro Asn Phe Asp Lys Val Lys Gly Val Ile 165 170 Cys Ser Val Gly Gly Gly Leu Tyr Asn Gly Ile Val Glu Gly Leu 185 Glu Asn His Lys Glu Ile Pro Val Leu Ala Ile Glu Thr Lys Gln Ala 205 200 195 Ala Thr Phe His Glu Ala Val Lys Glu Gly Lys Val Val His Leu Gln 210 215 220 Lys Val Gln Thr Leu Ala Thr Ser Leu Ala Ser Pro Tyr Leu Ser Ser 240 230 235 225 Lys Ala Leu Ala Asn Tyr Ile Glu Arg Pro Thr Val Leu Ala Glu Ile 245 250 Asp Asp Leu Asp Ala Val Lys Gly Val Val Asp Val Tyr Asp His Phe 260 265 Gly Tyr Met Val Glu Pro Ala Cys Gly Ala Ser Val Ala Ser Val Met 280 275

His Arg Gln Asp Leu Leu Asn Lys Phe Gly Thr Leu Ser Pro Asp Asp 290 295 300

Ile Ile Ile Val Val Ile Cys Gly Gly Ser Ala Ile Asn Lys Tyr Ile 305 310 315 320

Ile Asp Glu Tyr Arg Ser Leu Leu Glu Lys Asp Ser 325 330

<210> 377

<211> 2564

<212> DNA

<213> Candida albicans

<400> 377

ttttgaaaa cagagaatga atgaacaatt gatgaccata gaaatgaagt gagaacacat 60 aaatctgcga cacttcacgt gataacaaca aagtgacatg aacaacaaat gtcgtttgta 120 taatttgcga catttgttgt tcaagccaaa aaaagaaaga cagaaacaga aaaaagaaaa 180 atctcaaagt tgccacgtaa gcacaattaa tctttttatg gttgtaacta tttctagtat 240 atctacgta ataggtgagg tcctatatgc agtacacaca ggttttttc acagatgttg 300 acacagtgtt gaaaattatt cacgttgatt tataaatata caacttacct tccacggttt 360

```
tttttttgat ttttgatttt ttgaattctt ctttctttt tttgtcattt atttattaaa 420
 acattetett gtaataettt atttgataaa ttttacaaga tttaattaga teaattagat 480
· tatagaatca tttcaatata atgagatcat cacaatcttc ttggttacca cgtattggtt 540
 tattatatgt ggcgttagtt atacttatac catttttggt ttcaccgaag cacgcatttg 600
 cggttgctgc agtgagcgat gatgaatcct cgacagataa ttatggtaca gtcattggta 660
 tagatttggg taccacttat tcctgtgttg gtgttatgaa aaacggtaag gttgaaattt 720
 tggccaacga tcaaggtaat agaatcactc catcatacgt gtcgttcaat ggcgatgaaa 780
 gattggttgg agacgctgcc aagaatcaag cttcctctaa tgtcaacaac actgttttcg 840
 atattaaaag attgattggt ttgaaatata atgatgacac tgtgcaaaag gaacttaaac 900
 atttgcctta caaaattgaa aataagggta acaaaccagt tgttaaagtt gaataccaag 960
 gtgaagaaaa aactttctcc cctgaagaaa tttcatctat ggttttgggt aaaatgaaga 1020
 gtattgctga agattacctt ggcaaaaaag ttactcatgc cgttgtcact gttccagctt 1080
 atttcaacga tgctcaaaga caagctacta aagatgccgg tactattgct ggtttgaacg 1140
 ttttgagaat tgtcaatgaa cctactgctg ccgctattgc ctatggatta gacaaaggcg 1200
 accaagaaaa acaaattatt gtttacgatt tgggtggtgg tacttttgat gtttctttat 1260
 tgtccattga aggtggtgtt ttcgaagtct tggctactgc tggtgatact cacttgggtg 1320
 gtgaagattt tgatttcaag attgtcagat acttggccaa acaattcaag aagaagcaca 1380
 atattgatat cactgccaat tctaaagcca tttccaaatt gaagagagaa gccgaaaagg 1440
 ccaagagaac tttatcttct caaatgagta ctagagttga aatcgactcc tttgttgatg 1500
 gtattgactt ttctgaaact ctttcaagag ccaagtttga agaattgaac attgctgctt 1560
 tcagaaagac tttgaaacca gttgaacaag tgttgaagga tggtggtgtc aagaaatccg 1620
 atattgatga tattgttttg gttggtggtt ccaccagaat tccaaaagtt caagaattat 1680
 tggaaggatt ctttgatggt aaaaaggctt ctaaaggtat taacccagat gaagctgttg 1740
 cttatggtgc cgctgttcaa gcaggtgttt tgagtggtga agaaggtgtt gatgacattg 1800
 ttttgttgga tgttaaccca ttgactttag gtattgaaac ttctggtggg gttatgacca 1860
 ctttgatcaa gagaaacact gccatcccaa ccaagaaatc tcaaatcttc tcaactgctg 1920
 ctgataacca accaactgtt ttgatccaag tctatgaagg tgaaagaacc atggctaaag 1980
 acaacaacag attgggcaaa ttcgaattga ctggtattcc accagctcca agaggtgtcc 2040
 cacaaattga agtcactttc tcattggatg ccaatggtat cttgaaagtt gaagctgctg 2100
 ataagggaac tggtaaatct gaatccatta ctatcaccaa cgaaaagggt agattatcca 2160
 aggatgaaat tgatagaatg gttgaagaag ctgaaaaata cgctcaacaa gatcaagaat 2220
 tgaaagaaaa gattgaagct agaaactcat tagaaaacta tgctcatgtc ttgagaggtc 2280
 aattgagtga tacttctgaa accggtttag gttctaaatt ggatgacgat gacaaggaaa 2340
 ctttggatga cgctatcaag gaaactttag aatttattga agataacttt gatactgcta 2400
 ctgctgaaga atttgaagaa caaaaacaaa aattaattga cgttgctaac ccaatcacag 2460
 caaaattata cggtggagct gctggtgaag gtgctggtgg cgctggggat gccaaattcg 2520
                                                                   2564
 gtgatgatga ttcagatgat gaattcgatc acgatgaatt gtag
```

Val	Ala	Leu	Val	Ile	Leu	Ile	Pro	Phe	Leu	Val	Ser	Pro	Lys	His	Ala
			20					25					30		

Phe Ala Val Ala Ala Val Ser Asp Asp Glu Ser Ser Thr Asp Asn Tyr 35 40 45

Gly Thr Val Ile Gly Ile Asp Leu Gly Thr Thr Tyr Ser Cys Val Gly 50 55 60

Val Met Lys Asn Gly Lys Val Glu Ile Leu Ala Asn Asp Gln Gly Asn 65 70 75 80

Arg Ile Thr Pro Ser Tyr Val Ser Phe Asn Gly Asp Glu Arg Leu Val 85 90 95

Gly Asp Ala Ala Lys Asn Gln Ala Ser Ser Asn Val Asn Asn Thr Val
100 105 110

Phe Asp Ile Lys Arg Leu Ile Gly Leu Lys Tyr Asn Asp Asp Thr Val 115 120 125

Gln Lys Glu Leu Lys His Leu Pro Tyr Lys Ile Glu Asn Lys Gly Asn 130 135 140

Lys Pro Val Val Lys Val Glu Tyr Gln Gly Glu Glu Lys Thr Phe Ser 145 150 155 160

Pro Glu Glu Ile Ser Ser Met Val Leu Gly Lys Met Lys Ser Ile Ala 165 170 175

Glu Asp Tyr Leu Gly Lys Lys Val Thr His Ala Val Val Thr Val Pro 180 185 190

Ala Tyr Phe Asn Asp Ala Gln Arg Gln Ala Thr Lys Asp Ala Gly Thr 195 200 205

Ile Ala Gly Leu Asn Val Leu Arg Ile Val Asn Glu Pro Thr Ala Ala 210 215 220

Ala Ile Ala Tyr Gly Leu Asp Lys Gly Asp Gln Glu Lys Gln Ile Ile 225 230 235 240

Val Tyr Asp Leu Gly Gly Gly Thr Phe Asp Val Ser Leu Leu Ser Ile 245 250 255

Glu Gly Gly Val Phe Glu Val Leu Ala Thr Ala Gly Asp Thr His Leu 260 265 270

Gly Glu Asp Phe Asp Phe Lys Ile Val Arg Tyr Leu Ala Lys Gln 275 280 285

Phe Lys Lys Lys His Asn Ile Asp Ile Thr Ala Asn Ser Lys Ala Ile 290 295 300

Ser Lys Leu Lys Arg Glu Ala Glu Lys Ala Lys Arg Thr Leu Ser Ser 305 310 315 320

Gln Met Ser Thr Arg Val Glu Ile Asp Ser Phe Val Asp Gly Ile Asp 325 330 335

Phe Ser Glu Thr Leu Ser Arg Ala Lys Phe Glu Glu Leu Asn Ile Ala 340 345 350

Ala Phe Arg Lys Thr Leu Lys Pro Val Glu Gln Val Leu Lys Asp Gly 355 360 365

Gly Val Lys Lys Ser Asp Ile Asp Asp Ile Val Leu Val Gly Gly Ser 370 375 380

Thr Arg Ile Pro Lys Val Gln Glu Leu Leu Glu Gly Phe Phe Asp Gly 385 390 395 400

Lys Lys Ala Ser Lys Gly Ile Asn Pro Asp Glu Ala Val Ala Tyr Gly
405 410 415

Ala Ala Val Gln Ala Gly Val Leu Ser Gly Glu Glu Gly Val Asp Asp 420 425 430

Ile Val Leu Leu Asp Val Asn Pro Leu Thr Leu Gly Ile Glu Thr Ser
435 440 445

Gly Gly Val Met Thr Thr Leu Ile Lys Arg Asn Thr Ala Ile Pro Thr 450 455 460

Lys Lys Ser Gln Ile Phe Ser Thr Ala Ala Asp Asn Gln Pro Thr Val 465 470 475 480

Leu Ile Gln Val Tyr Glu Gly Glu Arg Thr Met Ala Lys Asp Asn Asn 485 490 495

Arg Leu Gly Lys Phe Glu Leu Thr Gly Ile Pro Pro Ala Pro Arg Gly 500 505 510

Val Pro Gln Ile Glu Val Thr Phe Ser Leu Asp Ala Asn Gly Ile Leu
· 515 520 525

 Lys
 Val
 Glu
 Ala
 Ala
 Asp
 Lys
 Gly
 Thr
 Gly
 Lys
 Ser
 Glu
 Ser
 Ile
 Thr

 530
 535
 535
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 540
 550
 550
 555
 555
 555
 560
 560
 560
 560
 555
 555
 560
 560
 560
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 575
 5

Gly Gln Leu Ser Asp Thr Ser Glu Thr Gly Leu Gly Ser Lys Leu Asp 595 600 605

Asp Asp Asp Lys Glu Thr Leu Asp Asp Ala Ile Lys Glu Thr Leu Glu 610 620

Phe Ile Glu Asp Asn Phe Asp Thr Ala Thr Ala Glu Glu Phe Glu Glu 625 630 635 640

Gln Lys Gln Lys Leu Ile Asp Val Ala Asn Pro Ile Thr Ala Lys Leu 645 650 655

Tyr Gly Gly Ala Ala Gly Glu Gly Ala Gly Gly Ala Gly Asp Ala Lys
660 . 665 670

Phe Gly Asp Asp Ser Asp Asp Glu Phe Asp His Asp Glu Leu 675 680 685

<210> 379

<211> 1346

<212> DNA

<213> Candida albicans

<400> 379

gccccatggt gacagaaaag tattgatcca ggtttctttg aaggtttgta atacgtttta 60 aacttgtgac ctaactcatt ttcataaaat tcttgtataa aattggtttc tttgaattgt 120 ttgtaatctc ttaagattgt ctcgtcttca gtgtttgcag tggagctgct attgccagtc 180 ggcggcataa cagcaggttc tggtatgtca tcttcatcta ctaagccgga caatcccaaa 240 gccgtctctt gttctttat tcttctaagg aatgctttat gtagttctga cattcttgcg 300 taaaaggaag tacagtatta aagctcgaaa ctcaattgca acaaaagtct taatttttc 360 ttctataagt aattgatctt ggtccccaa acaaaaagat acagaggaga aaggagggcc 420 ttaggctgac agaaaaaaa aatttagtac ctgccacac agtagctcca aacccatata 480 tgatttgata gagttgaagt atgctgacag atcttacttc acactttcag tacatggcta 540 taagcttgtt tgtcggttac aaagccttac taaataatga gacaccagtg tcatgtatag 600

```
tegtggatte aaaatcagat aaaatcatta geataggeta caattatace aaccaetete 660 taaatggtac acaacaegea gaatttattg etttacaaeg atttggggaa caaaaaetga 720 gtattgacta taatgatta attttgtatg tgacagtgga geettgtatt atgtgtgeat 780 ettattaceg teagttggg ateaaaaaag taatatttgg ttgtgggaat gatagatteg 840 gaggaaatgg taccattta teaatacata gtgacateae ettgeetaae geagettatt 900 eeagtategg gggtatatgt aggacagaag gaatecaaet attacgaaat tettatatte 960 aacaaaaatga gteggeaeca aatecaaaaa teaaaaagaa cacagatatt gagagtaagg 1020 aatacecaga gaateagtt tgtageattt eaaaggatga atttatagag tettaceggaa 1080 atgaaaggat geatattae gatgggaaga tetttgaaat eactecatta caaaacaagg 1140 gttatgatat aaaagaattg atategttgg atatgatga aaaagteea tettetagagg 1200 atgagttggg tggaacggt aattataaa aaccaatagg caaatataae agtaaaaaga 1320 gacaetttge aaacgatgaa gaatag
```

<210> 380

<211> 281

<212> PRT

<213> Candida albicans

<400> 380

Met Ser Thr Asp Leu Thr Ser His Phe Gln Tyr Met Ala Ile Ser Leu 1 5 10 15

Phe Val Gly Tyr Lys Ala Leu Leu Asn Asn Glu Thr Pro Val Ser Cys 20 25 30

Ile Val Val Asp Ser Lys Ser Asp Lys Ile Ile Ser Ile Gly Tyr Asn 35 40 45

Tyr Thr Asn His Ser Leu Asn Gly Thr Gln His Ala Glu Phe Ile Ala 50 55 60

Leu Gln Arg Phe Gly Glu Gln Lys Ser Ser Ile Asp Tyr Asn Asp Leu 65 70 75 80

Ile Leu Tyr Val Thr Val Glu Pro Cys Ile Met Cys Ala Ser Tyr Leu 85 90 95

Arg Gln Leu Gly Ile Lys Lys Val Ile Phe Gly Cys Gly Asn Asp Arg 100 105 110

Phe Gly Gly Asn Gly Thr Ile Leu Ser Ile His Ser Asp Ile Thr Leu 115 120 125

Pro Asn Ala Ala Tyr Ser Ser Ile Gly Gly Ile Cys Arg Thr Glu Gly
130 135 140

 Ile Gln Leu Leu Arg Asn Phe Tyr Ile Gln Gln Asn Glu Ser Ala Pro

 145
 150
 155
 160

 Asn Pro Lys Ile Lys Lys Asn Thr Asp Ile Glu Ser Lys Glu Tyr Pro
 165
 170
 175

 Glu Asn Gln Phe Cys Ser Ile Ser Lys Asp Glu Phe Ile Glu Phe Tyr
 185
 190

Gly Asn Glu Arg Val His Ile Tyr Asp Gly Lys Ile Phe Glu Ile Thr 195 200 205

Pro Leu Gln Asn Lys Gly Tyr Asp Ile Lys Glu Leu Ile Ser Leu Asp 210 215 220

Met Met Gln Lys Val Pro Phe Leu Glu Asp Glu Leu Gly Gln Ile Thr 225 230 235 240

Asp Glu Gln Ile Ile Glu Phe His Asn Leu Phe Phe Asn Ile Asn Asp 245 250 255

Asp Gly Thr Val Asn Tyr Lys Lys Pro Ile Gly Lys Tyr Asn Ser Lys 260 265 270

Lys Arg His Phe Ala Asn Asp Glu Glu 275 280

<210> 381 <211> 1504 <212> DNA <213> Candida albicans

<400> 381

<210> 382

<211> 334

<212> PRT

<213> Candida albicans

<400> 382

Met Leu Arg Phe Thr Arg Thr Thr Ala Trp Lys Leu Arg Ser Ile Pro 1 5 10 15

Ile Ala Thr Ile Gln Tyr Arg Gln Phe Thr Tyr Ser Thr Ile Cys Tyr
20 25 30

Gln Leu Lys Thr Leu Thr Pro Ser Leu Gly Ile Asn Asn Thr Ile Glu 35 40 45

Ser Asn Ile Pro Ser Glu Thr Asn Arg Leu Ala Lys Thr Gly Thr Arg
50 55 60

Phe Trp Lys Lys Gly Glu Val Lys Phe Asn Asn Glu Thr Gln Lys Tyr
65 70 75 80

Glu Ile Gln Leu Asp Gly Lys Thr Leu Arg Thr Pro Leu Gly Phe Pro 85 90 95

Leu Glu Leu Pro Ile Asn Lys Lys Gln Leu Ala Tyr Leu Ile Ala His
100 105 110

Glu Trp Thr His Leu Pro Asp Ile Lys Val Lys Ser Ser Thr Leu Pro 115 120 125

Leu Thr Ala Leu Ala Thr Arg Ala Ile Asp Leu Ser Gln Gln His Leu 130 135 140 Ser Asp Met Lys Thr Glu Lys Ala Glu Glu Met Leu Ala Leu Glu Asp 145 150 155 160

Ile Lys Leu Gln Met Leu Arg Tyr Leu Asp Thr Asp Thr Cys Leu Ile 165 170 175

Phe Ala Thr Asn Lys Glu Cys Asp Gly Lys Leu Arg Lys Arg Gln Glu 180 185 190

Glu Ile Tyr Arg Pro Leu Ile Asn Glu Phe Asn Glu Phe Phe Thr Ile 195 200 205

Tyr Ala His Asn Lys Asn Leu Ile Pro Arg Gln Lys Ser Ile Glu Leu 210 215 220

Lys Tyr Leu Asp Cys Glu Thr Asp Gly Leu Arg Gly Asn Lys Gln Asp 225 230 235 240

Glu Thr Thr Gln Leu Val Val Leu Asp Trp Leu Asn Gln Leu Pro Ile 245 250 255

Tyr Asp Leu Ile Ala Leu Glu Lys Thr Ile Leu Thr Thr Lys Ser Phe 260 265 270

Leu Cys Gly Ile Thr Leu Leu Arg Ser Asn Val Asn Asp Ile Glu Thr 275 280 285

Leu Lys Glu Leu Tyr Gln Phe Asn Lys Asn Ser Ile Asp Glu Asp Tyr 290 295 300

Tyr His Lys Thr Leu Glu Glu Leu Val Glu Leu Gly Asn Leu Glu Thr 305 310 315 320

Ile Tyr Gln Thr Glu Glu Trp Gly Glu Val Glu Asp Thr His
325 330

<210> 383

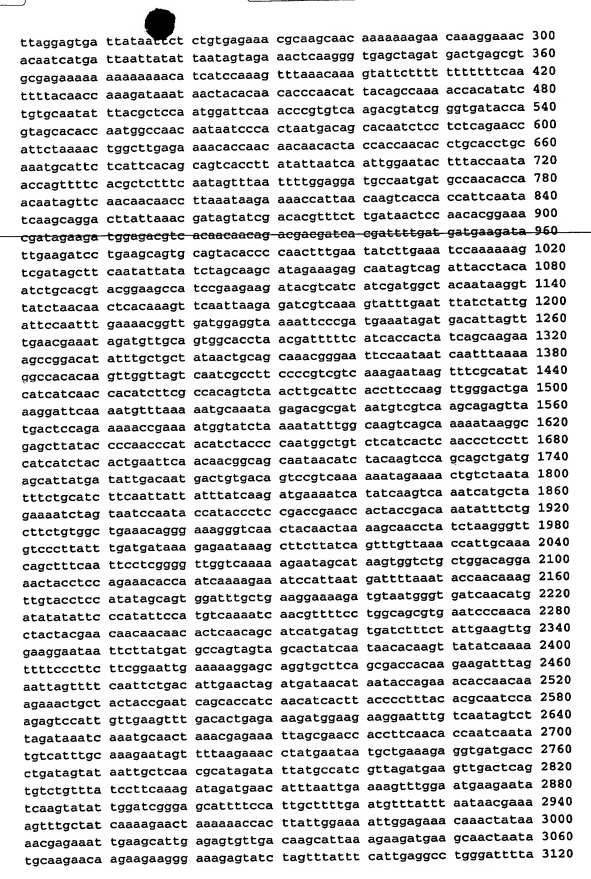
<211> 3689

<212> DNA

<213> Candida albicans

<400> 383

tatttatata taatctaaca acaagtacta tcccttaacc tattagctca atgtcattct 60 tcattaaaca agagatatgg taattgttga ttagacaaca gttgggtatc catttgtatt 120 ggaacgcttt ttttaaactg ctgttttatt gtaaaaatta ttgtcgtctt tcttcataac 180 attatttgta acagtctcta actgtattgt tgggctgaca gtagagtatt gcggtcattt 240



<210> 384

<211> 1062

<212> PRT

<213> Candida albicans

<400> 384

Met Asp Ser Asn Pro Cys Gln Asp Val Ser Gly Asp Thr Ser Ser Thr 1 5 10 15

Pro Met Ala Asn Asn Asn Pro Thr Asn Asp Ser Thr Ile Ser Ser Gln 20 25 30

Asn His Ser Lys Thr Gly Leu Arg Lys His Gln Gln His Tyr His
35 40 45

Gln His Ser His Ser Gln Met His Ser His Ser Gln Gln Ser Pro Tyr
50 55 60

Ile Asn Gln Leu Glu Tyr Phe Thr Asn Asn Gln Phe Ser Arg Ser Phe 65 70 75 80

Asn Ser Leu Ile Leu Glu Asp Ala Asn Asp Ala Asn Thr Asn Asn Ser 85 90 95

Ser Thr Thr Leu Asn Lys Lys Thr Ile Asn Lys Ser Pro Pro Phe 100 105 110

Asn Ile Lys Gln Asp Leu Leu Asn Asp Ser Ile Asp Thr Phe Leu Asp 115 120 125

Asn Ser Asn Thr Glu Thr Ile Glu Asp Gly Asp Val Thr Thr Asp 130 135 140

Asp Asp His Asp Phe Asp Asp Glu Asp Ile Glu Asp Pro Glu Ala Val 145 150 155 160

Gln Tyr Thr Pro Thr Leu Asn Ile Leu Lys Ser Lys Lys Val Asp Ser 165 170 Phe Asn Ile Ile Ser Ser Lys His Arg Lys Ser Asn Ser Gln Ile Thr . 185 180 Tyr Asn Ser His Val Arg Lys Pro Ser Glu Glu Asp Thr Ser Ser Ser 200 Met Ala Thr Ile Arg Leu Ser Asn Asn Ser Gln Ser Ser Ile Lys Arg 215 Ser Ser Lys Tyr Leu Asn Leu Ser Ile Asp Ser Asn Leu Lys Thr Val 225 230 240 Asp Gly Gly Lys Ile Pro Asp Glu Ile Asp Asp Ile Ser Leu Asn Glu 245 250 Ile Asp Val Ala Val Ala Pro Asn Asp Phe Ser Ser Pro Leu Ser Ala 260 265 Arg Lys Pro Asp Ile Phe Ala Ala Ile Thr Ala Ala Asn Gly Asn Ser 280 Asn Asn Gln Phe Lys Arg Pro His Lys Leu Val Ser Gln Ser Pro Ser 295 Pro Ser Ser Lys Asn Lys Phe Arg Ile Ser Ser Ser Thr Thr Ser Ser 310 315 Pro Gln Ser Asn Leu His Ser Pro Ser Lys Leu Gly Ser Lys Gly Phe 325 330 335 Lys Met Phe Lys Asn Ala Asn Arg Asp Ala Ile Met Ser Ser Ser Arg 340 345 Val Met Thr Pro Glu Lys Pro Lys Met Val Ser Lys Ile Phe Gly Lys 355 360 Ser Ala Lys Ile Arg Arg Ala Tyr Thr Pro Thr His Thr Ser Thr Pro 370 375 Met Ala Val Ser Ser Leu Asn Pro Pro Ser Ser Ser Thr Ser Asn Ser 390 395 Thr Thr Ala Ala Ile Thr Ser Thr Ser Pro Ala Ala Asp Glu His Tyr

461

410

Asp	Ile	Asp	Asn 420	Asp	Cys	Asp	Ser	Pro 425	Ser	Lys	Asn	Arg	Lys 430	Ser	Ser
Asn	Ile	Ser 435	Ala	Ser	Ser	Ile	Ile 440	Ile	Tyr	Gln	Asp	Glu 445	Asn	His	Ile
Lys	Ser 450	Asn	His	Ala	Arg	Lys 455	Ser	Ser	Asn	Pro	Ile 460	Pro	Tyr	Pro	Pro
Thr 465							Ile				Val	Ala	Glu	Thr	Gly 480
Lys	Gly	Ser	Thr	Thr 485	Thr	Lys	Ser	Asn	Leu 490	Ser	Lys	Gly	Cys	Pro 495	Leu
Phe	Asp	Asp	Lys 500	Glu	Asn	Lys	Ala	Ser 505	Tyr	Gln	Phe	Val	Lys 510	Pro	Leu
Gln	Thr	Ala 515	Phe	Asn	Ser	Ser	Gly 520	Leu	Val	Lys	Lys	Asn 525	Ser	Ile	Ser
Gly	Ser 530	Ser	Asp	Arg	Lys	Leu 535	Pro	Pro	Glu	Thr	Pro 540	Ile	Lys	Arg	Asn
Pro 545	Leu	Met	Ile	Leu	Asn 550	Thr	Asn	Lys	Val	Val 555	Pro	Pro	Tyr	Ser	Ser 560
Gly	Phe	Ala	Glu	Gly 565	Lys	Asp	Val	Met	Gly 570	Asp	Gln	His	Asp	Ile 575	Tyr
Ser	His	Ile	Pro 580	Cys	Gln	Asn	Gln	Arg 585	Phe	Pro	Gly	Ser	Val 590	Asn	Pro
Asn	Thr	Thr 595	Thr	Asn	Asn	Asn	Asn 600	Thr	Gln	Gln	His	His 605	Asp	Ser	Asp
Leu	Ser 610	Ile	Glu	Val	Gly	Arg 615	Asn	Asn	Ser	Tyr	Asp 620	Ala	Ser	Ser	Ser
Thr 625	Ile	Asn	Asn	Thr	Ser 630	Tyr	Ile	Lys	Ile	Phe 635	Pro	Ser	Ser	Glu	Leu 640
Lys	Lys	Glu	Gln	Val 645	Leu	Gln	Arg	Pro	Gln 650	Glu	Asp	Leu	Glu	Leu 655	Val
Phe	Asn	Ser	Asp 660	Ile	Glu	Leu	Asp	Asp 665	Asn	Ile	Ile	Pro	Glu 670	Thr	Pro

Thr Lys Lys Ser Leu Leu Pro Asn Gln His His Gln His His Leu Pro 675

Leu Tyr Thr Gln Ser Lys Ser Pro Leu Leu Lys Phe Asp Thr Glu Lys 690

Asp Gly Arg Arg Asn Leu Ser Ile Val Leu Asp Lys Ser Asn Ala Thr 705

Lys Arg Glu Ile Ser Glu Pro Pro Ser Thr Pro Ile Asn Met Ser Phe 735

Ala Lys Asn Ser Phe Lys Lys Pro Met Asn Asn Ala Glu Arg Gly Asp 740 745 750

Asp Pro Asp Ser Ile Ile Ala Gln Arg Ile Asp Ile Met Pro Ser Leu 755 760 765

Asp Glu Val Asp Ser Val Ser Val Tyr Pro Ser Lys Ile Asp Glu His
770 775 780

Leu Ile Glu Lys Phe Gly Met Lys Asn Ile Lys Tyr Ile Gly Ser Gly 785 790 795 800

Ala Phe Ser Ile Ala Phe Glu Cys Leu Phe Asn Asn Glu Lys Phe Ala 805 810 815

Ile Lys Arg Thr Lys Lys Pro Leu Ile Gly Lys Leu Glu Lys Gln Thr 820 825 830

Ile Lys Arg Glu Ile Glu Ala Leu Arg Val Leu Thr Ser Ile Lys Glu 835 840 845

Asp Glu Ala Thr Asn Met Gln Glu Glu Glu Glu Gly Lys Glu Tyr Leu 850 855 860

Val Tyr Phe Ile Glu Ala Trp Asp Phe Asn Asn Tyr Tyr Tyr Ile Met 865 870 875 880

Thr Glu Phe Cys Glu Gly Gly Thr Leu Phe Asp Phe Leu Glu Glu Asn 885 890 895

Lys His Tyr Lys Ile Asp Glu Phe Arg Ile Trp Lys Ile Leu Ile Glu 900 905 910

Ile Leu Asn Gly Leu Lys Phe Ile His Ser Lys Asn Tyr Leu His Leu 915 920 925

Ile Leu Glu Ile Ala Ala Asn Ile Ile Leu Pro Asp Asn Gly Thr Pro 995 1000 1005

Trp Arg Lys Leu Arg Ser Gly Asp Leu Ser Asp Ala Gly Arg Leu Ser 1010 1015 1020

Ser Asp Asn Ile Ser Met Phe Leu Gln His Asn Pro Asn Thr Asn Ser 1025 1030 1035 1040

Asn Ile Ser Gly Ser Gly Ser Arg Ser Gly Ser Gly Ser Thr Gly Gly
1045 1050 1055

Asn Gly Ser Ala Gly Asp 1060

<210> 385 <211> 887 <212> DNA <213> Candida albicans

<400> 385

agtcaagaat tttatgatga ctttggagga tggtctgatt aagacttgtc ttttaccagt 60 tttttcagca ttgttgatgg cgtttaaagc atcagctaag acggaggttc tagtcatggt 120 tgaattgtct gttgataaa agaaaccgtg aaaaggaagt attgtgaaaa atcgattgat 180 atttttttt ttcctccttc ctcactgtaa cagtagtaaa cacactagtt acaactgatg 240 acctgcatat tataaatctt tctgaaaaaa ttttttccc tgtatttttg taattctttc 300 gctcttctc actcactcac acttattaat gaatgaaagg tttggtgtct acaaactcca 360 ctaacaaaat ctcactcctg tgcctaaaca cacacagacc cacacgcaaa cctttctctc 420 agaaacagaa aaaaaaaatt tcaagcaaaa atttttcca tctagattt ctttctcag 480 aatatcaagt aactttaaag atgcctgtat gtacaataac agtgttataa tggatgtcttt 540 cagtcttta ttgggtgatt attataataa atggcagatg agaaatatat tggataataa 600 accttgcaaa tgaaatcagt tgaccatttt gaaataatgg accattagca atacccagat 660 tgctggaaca aagacgagag aactagacaa ctcagaactaa acaaaagtta gctaaggctc 780

02-07-1999

```
aaaagcaaaa cagaccattg ccacaatgga tcagattgag aactgacaac aaaatcagat 840 acaatgctaa aagaagacac tggagaagaa ctaagttggg tatctaa 887
```

<210> 386

<211> 51

<212> PRT

<213> Candida albicans

<400> 386

Met Pro Ser Gln Lys Ser Phe Arg Thr Lys Gln Lys Leu Ala Lys Ala

Gln Lys Gln Asn Arg Pro Leu Pro Gln Trp Ile Arg Leu Arg Thr Asp 20 25 30

Asn Lys Ile Arg Tyr Asn Ala Lys Arg Arg His Trp Arg Arg Thr Lys
35 40 45

Leu Gly Ile 50

<210> 387

<211> 893

<212> DNA

<213> Candida albicans

<400> 387

gatttcattt gcaaggttta ttatccaata tatttctcat ctgccattta ttataataat 60 cacccaataa aagactgaaa gacatcatta taacactgtt attgtacata caggcatctt 120 taaagttact tgatattctg aagaaagaaa atctagatgg aaaaaatttt tgcttgaaat 180 ttttttttc tgtttctgag agaaaggttt gcgtgtgggt ctgtgtgtgt ttaggcacag 240 gagtgagatt ttgttagtgg agtttgtaga caccaaacct ttcattcatt aataagtgtg 300 agtgagtgag aaagagcgaa agaattacaa aaatacaggg aaaaaaattt tttcagaaag 360 atttataata tgcaggtcat cagttgtaac tagtgtgttt actactgtta cagtgaggaa 420 ggaggaaaaa aaaaaatatc aatcgatttt tcacaatact tccttttcac ggtttctttt 480 tatcaacaga caattcaacc atgactagaa cctccgtctt agctgatgct ttaaacgcca 540. tcaacaatgc tgaaaaaact ggtaaaagac aagtcttaat cagaccatcc tccaaagtca 600 tcataaaatt cttgactgtc atgcaaaaac acggttacat tggtgaattc gaatacattg 660 atgatcacag atccggtaaa attgttgttc aattaaatgg tagattaaac aaatgtggtg 720 tcattcaacc aagattcaac gtcaaaatca acgacattga aagatggact gacaacttgt 780 tgccagctag acaattcggt tacgttatct taaccacttc tgctggtatc atggaccacg 840 aagaagctag aagaaagcac gtttctggta aaatcttagg tttcgtttac tag 893

<210> 388

Val Ile Ile Lys Phe Leu Thr Val Met Gln Lys His Gly Tyr Ile Gly
35 40 45

Glu Phe Glu Tyr Ile Asp Asp His Arg Ser Gly Lys Ile Val Val Gln 50 55 60

Leu Asn Gly Arg Leu Asn Lys Cys Gly Val Ile Gln Pro Arg Phe Asn 65 70 75 80

Val Lys Ile Asn Asp Ile Glu Arg Trp Thr Asp Asn Leu Leu Pro Ala 85 90 95

Arg Gln Phe Gly Tyr Val Ile Leu Thr Thr Ser Ala Gly Ile Met Asp 100 105 110

His Glu Glu Ala Arg Arg Lys His Val Ser Gly Lys Ile Leu Gly Phe 115 120 125

Val Tyr 130

<210> 389 <211> 4619 <212> DNA <213> Candida albicans

<400>-389
caacgacaac aacaataaga agaatettt ccagatattg taaaatectt atattaaatg 60
ttaatagaag cctaaaaatt gcttgaattg atagatttgc tgattetttt gctagaaaca 120
cgcgttatat tacttatect ttetgtaatt tettgatete tgtttgaate aacaacgeet 180
ttgttttete tttetgeete cetetgeeae eecagttta tttgtttget tgtegaegtg 240
ctgccaaaaa aaaaaattga gttetetete tetttttae aacagagaag agacaaagaa 300
aaaaaaaaae atcaaaatta agatteaage ttttttttt agtttttta acaacaaaat 360
tgaaattaca atcettgaat ttacaacte atatteaget etaaatacta ataattataa 420
caataattaa ttgaattatt etataaacca etgatatttg atttatett tttattttg 480

tttgttccca	taattgtgtc	atgcctgaca	atatagaaga	tcgatccgag	ataccttctg	540
		acaaatgaaa				
		caaggtgaat				
		caattaatca				
		atccctcaaa				
		aaagatcaac				
		cccgagaatg				
		ggtgaatggt				
		accaaagaaa				
		aagacgcaac				
		catggtcatc				
		ttaatggatg				
		aggatttggt				
		tttatgtttg				
		aaagatcacg				
		caaacagaat				
		tcacagggtg				
		gctttacaat				
		aaaaaagaat				
		ggttcacttt				
		gaattcaaat				
		tctcaagagt				
		caaaaaccat				
		atcaccaaac				
		attgatttat				
		tccataactt				
		tatcacacat				
		aagttggaag				
		aatttcttga				
		tatagtgact				
		agagatacag				
		gtgccagtgt				
		ggaatcccat				
		agaaagaaat				
		gaaaaaataa				
		ttccctgcat				
		agcgacaacg				
		ttggctaacc				
-		agccctaaat				
_		aaacaaacag				
		cctttatcgg				
		aagatggatg				
		ngagaagcga				
		tttgtattaa				
		gaaacggtac				
		gatgaagaaa				
_		ccgttgcctc				
-		atggaaagta				
Cittyaatyt		acygudayta	accegaaag	Juliususus	2	

```
ctggtacaac attgatatcg aaggacacag ttttgttatg tgattgggat aaggaaattt 3420
atcaaaaatg ctttggtgat aaagaattac aagcatggga aaacatatcg aatttaccga 3480
atccagaatt ggagaaaaat agagctcatt ttgaaagaca aagaaaagct aaaattacat 3540
tatctgattg tcttaagagt ttcagtaccc ctgaaatttt aggtgaacat gatttatggt 3600
attgtccacg ttgtactgaa cataaacgtg ccacaaagac aatccaactt tggtcaacgg 3660
gtgatatcct cactattcat ttgaaaagat ttcatagtgc tcgtgcattt agtgataaga 3720
ttgatgtttt ggttgatttc ccaattgaag gtttagatat aagttcgtat gttgccaata 3780
ctgatttgac acctgaagat tgtttatacg acttgattgc cgttgataat cattatggtg 3840
ggttaggagg tggtcattac actgcctcgg taaagaattt cagagatgat aaatggtatt 3900
attttaatga tagtcgagtc actgaaatta ataatcctca agaagtcgta gctaattctg 3960
cgtacctttt attttaccgt cgaagaagtt cgaaaggagc tggtattttg ggaggagaaa 4020
actttatcga cttgcttcaa aaaggtcgag aggaatactc tgagagtttg caaaagaaaa 4080
gattggttct tcaaaatgtt ggccaaatag tcaatacgta tgccaaaatt gaacaagata 4140
taattgataa agaaacagag aaacagaaag aggaacaaga acaggaacag gaacaggaac 4200
aggaacagga acaagagcaa gagcaagagc cagttcaaga gccagatcaa gaacaagagc 4260
cagatcaaga gccagatcaa gatcaagatc aagagccaga tcaagagcca gatcaagatc 4320
aagagcagaa tgaaacaata aaaaaatcta gaccattcga tgaactcaaa ccatcaacta 4380
gtgaaacaaa taaccaacaa caaacaactc agttcaactt tgatgatgaa gataatgatt 4440
acgattatga agcagaagta gaagattcca atattcgcaa acaaagatta ctttcaaaag 4500
aaaataacag caataaattg gtgcatatta aaagcaatgg tcgccaagaa gtcacttcat 4560
caccagtacc aattgaaact gatggtgaca ctgatgtaac tgattccaat tcaacatag 4619
```

<210> 390

<211> 1372

<212> PRT

<213> Candida albicans

<400> 390

Met Pro Asp Asn Ile Glu Asp Arg Ser Glu Ile Pro Ser Asp Ala Lys

1 5 10 15

Glu Ile Val Thr Thr Asn Glu Ile Glu Ala Thr Asp Ser Glu His Thr
20 25 30

Thr Asn Val Asp Asn Glu Leu Pro Gln Gly Glu Ser Asn Glu Gln Thr
35 40 45

Gly Asp Asp Ser Asn Asp Asn Leu Ala Ser Lys Arg Gln Leu Ile Asn 50 55 60

Asp Leu Leu His Asn Asp His Phe Glu Glu Gly Thr Glu Arg Tyr Ile 65 70 75 80

Ile Pro Gln Asn Phe Leu His Glu Phe Leu Asn Leu Pro Ile Asp Asn 85 90 95

Phe Ser Asp Leu Lys Asp Gln Leu Gly Pro Ile Asp Phe His Ser Leu

100 105 110

Leu Asn Glu Gln Gly Asn Leu Tyr Pro Glu Asn Glu Glu Pro Val Thr
115 120 125

Phe Cys His Val Ser Pro Glu Val Phe Gln His Leu Gly Glu Trp Phe 130 135 140

Gly Ile Leu Gly Gln Pro Ile Ile Arg Ala Ile Ile Ile Asn Pro Asp 145 150 155 160

Thr Lys Glu Lys Gln Ile Glu Arg Phe Pro Pro Leu Phe Trp Val His 165 170 175

Gln Leu Gly Lys Lys Thr Gln Pro Thr Tyr Leu Arg His Arg His Asn 180 185 190

Gly Ser Asn His Asn His His His Gly His His Asp Ser Pro Ile 195 200 205

Pro Val Leu Leu Ser Lys Thr Ser Thr Phe His Arg Leu Met Asp Val 210 215 220

Ile Arg Tyr Asn Val Leu Lys Ala Pro Arg Lys Ser Thr Lys Asp Phe 225 230 235 240

Arg Ile Trp Phe Ile Val Pro Gln Asp Lys Gly Leu Gln Tyr Leu Ile 245 250 255

Ser Ile Gln Thr Phe Met Phe Asp Ile Ser Lys Lys Thr Leu Val Ser 260 265 270

Pro Asn Met Leu Glu Asp Ala Leu Lys Asp His Gly Ile Val Ala Ser 275 280 285

Ser Tyr Asn Ile Met Val Glu Ala Lys Glu Lys His Gln Thr Glu Phe 290 295 300

Pro Ile Asp Gln Phe Ile Leu Ser His Ser Asn Ala Tyr Glu Glu Val 305 310 315 320

Ser Gln Gly Gly His Leu Gly Leu Ser Asn Met Gly Asn Thr Cys 325 330 335

Tyr Met Asn Ser Ala Leu Gln Cys Leu Leu His Val Pro Glu Ile Asn 340 345 350

Tyr Tyr Phe Phe Tyr Asn Ile Tyr Lys Lys Glu Leu Asn Phe Asp Asn

355

360

365

Pro Leu Gly Tyr His Gly Asp Val Ala Asn Ala Phe Gly Ser Leu Leu 370 375 380

Lys Gln Ala Phe Asp His Val Lys Asn Ser Ser Ser Ile Ser Pro Arg 385 390 395 400

Glu Phe Lys Ser Thr Ile Gly Arg Tyr Ser Ser Met Phe Ser Gly Tyr
405 410 415

Leu Gln Gln Asp Ser Gln Glu Leu Leu Ser Trp Leu Leu Asp Ala Leu 420 425 430

His Glu Asp Leu Asn Arg Ile His Gln Lys Pro Tyr Cys Glu Lys Pro 435 440 445

Glu Leu Lys Asp Asp Glu Ile Asp Asp Pro Gln Ala Ile Thr Lys Leu 450 455 460

Ala Asn Thr Cys Trp Asn Gln His Lys Ala Arg Asn Asp Ser Val Ile 465 470 475 480

Ile Asp Leu Phe Thr Gly Leu Tyr Gln Ser Thr Leu Ile Cys Pro Asp
485 490 495

Cys Gly Lys Lys Ser Ile Thr Phe Asp Pro Phe Asn Asp Leu Thr Leu 500 505 510

Pro Leu Pro Ile Ser Lys Lys Trp Tyr His Thr Phe Thr Ile Val Asp 515 520 525

Leu Ser Asn Gln Gly Val Ile Pro Glu Arg Ile Met Lys Leu Glu Val 530 540

Glu Leu Asn Lys Thr Ser Asn Phe Asp Asp Leu Leu Ser Tyr Leu Ser 545 550 555 560

Asn Phe Leu Asn Val Pro Ser Thr Glu Leu Phe Ala Tyr Glu Ile Phe
565 570 575

Gln Asn Ala Ile Tyr Ser Asp Phe Gln Leu Asp Tyr Thr Lys Asn Lys 580 585 590

Phe Leu Pro Ile Ser Asp Ile Ile Arg Asp Thr Asp Asp Val Ile Val 595 600 605

Tyr Ile Val Pro His Asn Pro Ala Val Asp Ile Ile Val Pro Val Phe

Asn Ala Val Glu Asp Ala Asp Ser Ser Tyr Gln Met Val Asn Phe Phe Gly Ile Pro Leu Phe Val Val Met Asn Lys Glu Val Asp Val Asn Ser Phe Gly Phe Ile Arg Lys Lys Leu Leu Glu Thr Val Ser Leu Leu Ser Lys Ile Asp Leu Val Asp Glu Tyr Glu Lys Ile Lys Arg Ser Asn Glu Asp Tyr Val Glu Lys Val Phe Tyr Lys Lys Ser Asp Phe Pro Ala Leu Ser Gln Pro Leu Glu Thr Ser Asp Cys Glu Lys Asn Asn Asn Asn Thr Ser Asp Asn Asp Asp Asp Glu Asp Ala Asp Asn Asp Glu Gly Tyr Asp Ser Glu Val Ser Leu Ala Asn Pro Tyr Leu Gly Ala Asn Phe Gly Phe Lys Ile Met Tyr Val His Asp Tyr Ser Pro Lys Leu Asn Ser Asn Leu Arg Ser Arg Tyr Asn His Asp Gln Thr Thr Lys Phe Lys Gln Thr Glu Arg Val Ile Asn Val Pro Thr His Lys Pro Thr Phe Ser Asp Phe Lys Pro Leu Ser Asp Gln Leu Ser Glu Ser Lys Arg Asn Tyr Tyr Phe Tyr Pro Asp Tyr Lys Lys Met Asp Asp Glu Met Asp Gln Leu Val Glu Glu .... 830 Val Asn Gln Asn Leu Ala Glu Gln Xaa Glu Ala Arg Ser Ser Gly Ser

Glu Asn Ser Ser Arg Ala Ser Glu Glu Gln Asp Gly Phe Val Leu Ile

Asn Lys Glu Asp Thr Leu Lys Gln Gln Ser Thr Val Pro Ala Ala Ala

875 880 870 865 Glu Thr Val Pro Pro Pro Leu Pro Val Arg Asn Asn Thr Gly Val His 885 890 Ile Pro Ser Ser Asp Glu Glu Thr Glu Ser Glu Ala Asn Leu Gly Ser 910 905 900 Leu Phe Asp Ser Thr Ser Asn Leu Pro Leu Pro Pro Pro Ser Thr Tyr 920 915 Ser Glu Ser Thr Lys Pro Ser Asn Val Asn Ser Pro Met Glu Ser Asn 935 940 930 Phe Glu Ser Ser Ser Ala Asp Leu Asn Ser Gly Thr Thr Leu Ile Ser 955 950 945 Lys Asp Thr Val Leu Cys Asp Trp Asp Lys Glu Ile Tyr Gln Lys 970 965 Cys Phe Gly Asp Lys Glu Leu Gln Ala Trp Glu Asn Ile Ser Asn Leu 985 980 Pro Asn Pro Glu Leu Glu Lys Asn Arg Ala His Phe Glu Arg Gln Arg 995 1000 Lys Ala Lys Ile Thr Leu Ser Asp Cys Leu Lys Ser Phe Ser Thr Pro 1015 1020 1010 Glu Ile Leu Gly Glu His Asp Leu Trp Tyr Cys Pro Arg Cys Thr Glu 1035 1025 1030

His Lys Arg Ala Thr Lys Thr Ile Gln Leu Trp Ser Thr Gly Asp Ile 1045 1050 1055

Leu Thr Ile His Leu Lys Arg Phe His Ser Ala Arg Ala Phe Ser Asp 1060 1065 1070

Lys Ile Asp Val Leu Val Asp Phe Pro Ile Glu Gly Leu Asp Ile Ser 1075 1080 1085

Ser Tyr Val Ala Asn Thr Asp Leu Thr Pro Glu Asp Cys Leu Tyr Asp 1090 1095 1100

Leu Ile Ala Val Asp Asn His Tyr Gly Gly Leu Gly Gly His Tyr 1105 1110 1115 1120

Thr Ala Ser Val Lys Asn Phe Arg Asp Asp Lys Trp Tyr Tyr Phe Asn



1130

1135

Asp Ser Arg Val Thr Glu Ile Asn Asn Pro Gln Glu Val Val Ala Asn 1140 1145 1150

Ser Ala Tyr Leu Leu Phe Tyr Arg Arg Arg Ser Ser Lys Gly Ala Gly 1155 1160 1165

Ile Leu Gly Gly Glu Asn Phe Ile Asp Leu Leu Gln Lys Gly Arg Glu 1170 1175 1180

Glu Tyr Ser Glu Ser Leu Gln Lys Lys Arg Leu Val Leu Gln Asn Val 1185 1190 1195 1200

Gly Gln Ile Val Asn Thr Tyr Ala Lys Ile Glu Gln Asp Ile Ile Asp 1205 1210 1215

Lys Glu Thr Glu Lys Gln Lys Glu Glu Glu Glu Glu Glu Glu Glu Gln Glu Gln 1220 1230

Glu Gln Glu Gln Glu Gln Glu Gln Glu Pro Val Gln Glu Pro 1235 1240 1245

Asp Gln Glu Gln Glu Pro Asp Gln Glu Pro Asp Gln Asp Gln Asp Gln 1250 1260

Glu Pro Asp Gln Glu Pro Asp Gln Asp Gln Glu Gln Asn Glu Thr Ile 1265 1270 1275 1280

Lys Lys Ser Arg Pro Phe Asp Glu Leu Lys Pro Ser Thr Ser Glu Thr 1285 1290 1295

Asn Asn Gln Gln Gln Thr Thr Gln Phe Asn Phe Asp Asp Glu Asp Asn 1300 1305 1310

Asp Tyr Asp Tyr Glu Ala Glu Val Glu Asp Ser Asn Ile Arg Lys Gln 1315 1320 1325

Arg Leu Leu Ser Lys Glu Asn Asn Ser Asn Lys Leu Val His Ile Lys 1330 1335 1340

Ser Asn Gly Arg Gln Glu Val Thr Ser Ser Pro Val Pro Ile Glu Thr 1345 1350 1355 1360

Asp Gly Asp Thr Asp Val Thr Asp Ser Asn Ser Thr 1365 1370 <210> 391 <211> 2693

```
<212> DNA
<213> Candida albicans
<400> 391
ttcatttcct aaattcaaat gaattatcat caccaatgcc accatcattt tctattaact 60
atggtagtga atgggattta gaaatcattc aaactagttt agataatgaa aaagaatcag 120
aaactaaatc atttactggt gaattagaat atacttcaac ttcttccaat ggtgaacatg 180
acaccaccac cactgccact aaacatgaat tgatattaca acaaattttg aattctaatg 240
atgaatcata tattaatcct aaatcattaa catttgatcc attaaaaatt ttcactaaac 300
aattaattgg tgaattaatt aaaattaatc aattttacaa ttcaaaagaa tcggaaattt 360
tcaaaattta taataattta attcatgatt tacaaaatca aaatattaat attgatgatg 420
tatttaaatt cactcaagct tataattatt ccgatccaaa tataataaat actgatgatc 480
atcatcaata tcatttaaaa tcaactttat caagaacagt tactaatgct agtgtatttg 540
ataccattaa tcatattgat aatgattatg ataataataa taacaaccaa aagaataatt 600
atgatttgga gaaacaaaat aatactacag ttgcaattca tgatgatgat gattcagaag 660
acgatgaaga agaagaggaa gaagaaactc atagtcatga ttcagtatta cttaatcata 720
cacattttaa tgttaaacaa caattaaaaa tcacattaaa acgtaaagcc attacattat 780
tttgtaaaaa atttgataaa acttgtggtt attcaattaa acaagatttt attaatgaat 900
ttttacctca atattctcga gtatttgaaa atgataccat tgaagaatta gattataaat 960
tgaatcaaat tattaaaatt tatgcctttt tatcaaataa attaactact caatcaacaa 1020
ctaaagaaga tttggataat ataaaatttg aattaagatc ttatttacgt gatcatattg 1080.
ttgatttaga taattctgtg gttcaaaata ataaaatggg tgatgaaggt catataatta 1200
attcaatgat gaatttatcc atgaaaagaa ttaatttacc acaatgcctt aaaaaattga 1260
ttaaatatga tcatattgat attccacaat ttttattaac tactcaaatg cttaaaatta 1320
ttattattgt cattgttttc attatattat tagcagtgaa aacttttaat gatccagttc 1380
aaggtcgttg tttagcagta ttagttgctg ctgccatgct ttgggcttca.gaagcattac 1440
ctttatacac tacagcttta ttaatcccac ttttggttgt tacttgtaaa gtttgtaaaa 1500
ctccgggaac cgatgatcca atggatgcca ccaaggcatc acaatatatt tttgggacaa 1560
tgtggaattc cacaattatg atattaattg gtgggtttac attagctgct gcattatcaa 1620
aatataatct tgccaaaata ttatcatcat atattttagc attagcaggt acaaatccaa 1680
gaaatgtatt attggcaatc atgtgtgtat cattatttct ttccatgtgg atttctaatg 1740
ttgctgcccc cgttttatgt ttttcattaa ttcaaccagt tttaagaagt atccccacag 1800
attcccccgt tgctaaagca ttagtgttag ggatcgcttt ggcgtctgat gttgctggta 1860
tggcttcacc aattgcatct ccacaaaatg ttattgctct tgaatcaatg aatcctaatc 1920
caggitgggg gaaatggitt getgtggcat tacctgtggc aatcattagt ttaattitaa 1980
tttgggtgga attattcatg acgtttaaaa tcaataatgt taaaatcaaa caattcaaac 2040
caattaaaga aaaattaacc atgaaacaat ggtttgtatt tgccgtcact ataactacta 2100
ttcttttatg gtgtgttatg caaaaaattg atggaacatt tggtgaatca ggtataatca 2160
cttgtatccc aattgtatta tttttcggta ccggtttatt aaaagttgat gatttaaata 2220
attatccttg gtcaattgtt atgttagcca tgggtggtat tgcattaggg aaagccgtta 2280
cttcttcagg tttattgaaa actattgctt tagcattaca aaaacgaatt atgcattatg 2340
atgccattgt tgtattaatc atttttggag cattaatttt ggtggtagct acatttgtaa 2400
gtcatactgt atcagcactt attattatcc ccttggttaa agaagttgga gattcattac 2460
```

ctaaacctca tccattaatg cttattatgg gtgtagcttt aattgcttca ggggcaatgg 2520 gattaccaac ttcaggattc cctaatgtga cggcaattgg tatgagagat gaagttggta 2580 aaccttattt gacggttaat ttattatta ctagaggggt tccggcaagt ataattgttt 2640 atgtttgtat tatcaccatt ggttatggta ttatgtcatc attgaacttt taa 2693

<210> 392

<211> 896

<212> PRT

<213> Candida albicans

<400> 392

His Phe Leu Asn Ser Asn Glu Leu Ser Ser Pro Met Pro Pro Ser Phe
1 5 10 15

Ser Ile Asn Tyr Gly Ser Glu Trp Asp Leu Glu Ile Ile Gln Thr Ser
20 25 30

Leu Asp Asn Glu Lys Glu Ser Glu Thr Lys Ser Phe Thr Gly Glu Leu 35 40 45

Glu Tyr Thr Ser Thr Ser Ser Asn Gly Glu His Asp Thr Thr Thr 50 55 60

Ala Thr Lys His Glu Leu Ile Leu Gln Gln Ile Leu Asn Ser Asn Asp 65 70 75 80

Glu Ser Tyr Ile Asn Pro Lys Ser Leu Thr Phe Asp Pro Leu Lys Ile 85 90 95

Phe Thr Lys Gln Leu Ile Gly Glu Leu Ile Lys Ile Asn Gln Phe Tyr 100 105 110

Asn Ser Lys Glu Ser Glu Ile Phe Lys Ile Tyr Asn Asn Leu Ile His 115 120 125

Asp Leu Gln Asn Gln Asn Ile Asn Ile Asp Asp Val Phe Lys Phe Thr 130 135 140

Gln Ala Tyr Asn Tyr Ser Asp Pro Asn Ile Ile Asn Thr Asp Asp His 145 150 155 160

His Gln Tyr His Leu Lys Ser Thr Leu Ser Arg Thr Val Thr Asn Ala 165 170 175

Ser Val Phe Asp Thr Ile Asn His Ile Asp Asn Asp Tyr Asp Asn Asn 180 185 190 Asn Asn Asn Gln Lys Asn Asn Tyr Asp Leu Glu Lys Gln Asn Asn Thr 195 200 205

Thr Val Ala Ile His Asp Asp Asp Ser Glu Asp Asp Glu Glu Glu 210 215 220

Glu Glu Glu Glu Thr His Ser His Asp Ser Val Leu Leu Asn His Thr 225 230 235 240

His Phe Asn Val Lys Gln Gln Leu Lys Ile Thr Leu Lys Arg Lys Ala 245 250 255

Ile Thr Leu Phe Ile Asn Leu Ser Glu Leu Lys Ser Phe Ile Glu Leu 260 265 270

Asn Arg Ile Gly Phe Thr Lys Ile Cys Lys Lys Phe Asp Lys Thr Cys 275 280 285

Gly Tyr Ser Ile Lys Gln Asp Phe Ile Asn Glu Phe Leu Pro Gln Tyr 290 295 300

Ser Arg Val Phe Glu Asn Asp Thr Ile Glu Glu Leu Asp Tyr Lys Leu 305 310 315 320

Asn Gln Ile Ile Lys Ile Tyr Ala Phe Leu Ser Asn Lys Leu Thr Thr 325 330 335

Gln Ser Thr Thr Lys Glu Asp Leu Asp Asn Ile Lys Phe Glu Leu Arg 340 345 350

Ser Tyr Leu Arg Asp His Ile Val Phe Glu Arg Asn Thr Val Trp Lys 355 360 365

Asp Leu Leu Ser Leu Glu Lys Lys Ser Tyr Asn Ile Asp Leu Asp Asn 370 375 380

Ser Val Val Gln Asn Asn Lys Met Gly Asp Glu Gly His Ile Ile Asn 385 390 395 400

Ser Met Met Asn Leu Ser Met Lys Arg Ile Asn Leu Pro Gln Cys Leu 405 410 415

Lys Lys Leu Ile Lys Tyr Asp His Ile Asp Ile Pro Gln Phe Leu Leu 420 425 430

Thr Thr Gln Met Leu Lys Ile Ile Ile Ile Val Ile Val Phe Ile Ile 435 440 445

Leu Leu Ala Val Lys Thr Phe Asn Asp Pro Val Gln Gly Arg Cys Leu 450 455 460

Ala Val Leu Val Ala Ala Ala Met Leu Trp Ala Ser Glu Ala Leu Pro 465 470 475 480

Leu Tyr Thr Thr Ala Leu Leu Ile Pro Leu Leu Val Val Thr Cys Lys
485 490 495

Val Cys Lys Thr Pro Gly Thr Asp Asp Pro Met Asp Ala Thr Lys Ala 500 505 510

Ser Gln Tyr Ile Phe Gly Thr Met Trp Asn Ser Thr Ile Met Ile Leu 515 520 525

Ile Gly Gly Phe Thr Leu Ala Ala Ala Leu Ser Lys Tyr Asn Leu Ala 530 535 540

Lys Ile Leu Ser Ser Tyr Ile Leu Ala Leu Ala Gly Thr Asn Pro Arg 545 550 555 560

Asn Val Leu Leu Ala Ile Met Cys Val Ser Leu Phe Leu Ser Met Trp 565 570 575

Ile Ser Asn Val Ala Ala Pro Val Leu Cys Phe Ser Leu Ile Gln Pro 580 585 590

Val Leu Arg Ser Ile Pro Thr Asp Ser Pro Val Ala Lys Ala Leu Val 595 600 605

Leu Gly Ile Ala Leu Ala Ser Asp Val Ala Gly Met Ala Ser Pro Ile 610 615 620

Ala Ser Pro Gln Asn Val Ile Ala Leu Glu Ser Met Asn Pro Asn Pro 625 630 635 640

Gly Trp Gly Lys Trp Phe Ala Val Ala Leu Pro Val Ala Ile Ile Ser 645 650 655

Leu Ile Leu Ile Trp Val Glu Leu Phe Met Thr Phe Lys Ile Asn Asn 660 665 670

Val Lys Ile Lys Gln Phe Lys Pro Ile Lys Glu Lys Leu Thr Met Lys 675 680 685

Gln Trp Phe Val Phe Ala Val Thr Ile Thr Thr Ile Leu Leu Trp Cys 690 695 700

Val Met Gln Lys Ile Asp Gly Thr Phe Gly Glu Ser Gly Ile Ile Thr 705 710 715 720

Cys Ile Pro Ile Val Leu Phe Phe Gly Thr Gly Leu Leu Lys Val Asp
725 730 735

Asp Leu Asn Asn Tyr Pro Trp Ser Ile Val Met Leu Ala Met Gly Gly
740 745 750

Ile Ala Leu Gly Lys Ala Val Thr Ser Ser Gly Leu Leu Lys Thr Ile
755 760 765

Ala Leu Ala Leu Gln Lys Arg Ile Met His Tyr Asp Ala Ile Val Val 770 780

Leu Ile Ile Phe Gly Ala Leu Ile Leu Val Val Ala Thr Phe Val Ser 785 790 795 800

His Thr Val Ser Ala Leu Ile Ile Ile Pro Leu Val Lys Glu Val Gly 805 810 815

Asp Ser Leu Pro Lys Pro His Pro Leu Met Leu Ile Met Gly Val Ala 820 825 830

Leu Ile Ala Ser Gly Ala Met Gly Leu Pro Thr Ser Gly Phe Pro Asn 835 840 845

Val Thr Ala Ile Gly Met Arg Asp Glu Val Gly Lys Pro Tyr Leu Thr 850 855 860

Val Asn Leu Phe Ile Thr Arg Gly Val Pro Ala Ser Ile Ile Val Tyr 865 870 875 880

Val Cys Ile Ile Thr Ile Gly Tyr Gly Ile Met Ser Ser Leu Asn Phe 885 890 895

<210> 393

<211> 2279

<212> DNA

<213> Candida albicans

<400> 393

tataaagttt caagtataaa aaggcgttta aaattaattc ttgtgattta aataagttaa 60

```
attettttt tttttetttg ttagttttte getttateeg ttttaeceaa gaaaggagea 120
atactgtttc accectecce cetteccagt tecatggece gecececec teteaacagt 180
ttattattaa agtcacaata cattgaaaga tagaatagca agcaaggtgg aaaaaaattt 300
ttttttttat cacattcaat acttatatcc taaattgata aacaatagag taattgatac 360
tttcgaagga caataaaacg atatatattt atatattaga gtgaacaata gcgattgcca 420
atcacagaca acagataatt tatctttctc ctattcattt gcacaattaa atccaaaaaa 480
aaaaaagaat cgaattccat atgtcgcata agactcagag ccaattatct tcacaaatga 540
aaaacttgaa tactccacca atagacttca actcaacttc aagtaacaat accatgcctt 600
ctgaaccaaa tctgcaaccg caacaacaac aatcacaacc agaagcaaaa acggagccac 660
aaaccatacg ccctgctact tttacaacta gtggcaattc atcatcttcg tcgatatcta 720
ccttatcage agatateatt caaccactte atcaactact gataaataac aacaattcaa 780
ctgtgacgca accagcgcca caaagctcat cgtttcaacg ccgaaacaat ccacaacgtt 840
tcaatcggaa tcaactcaat gtatacactg acttcaatag tactacttca tctgcttcaa 900
gcattagtag ttcaccaaaa gatttcttca ccagagagcc accacggatc catagtaaat 960
tgatatgtga agagattgcc tctgccaata atcgagctgc taaagaggtt ttatcacgtt 1020
tatctactga tgaattgcgt tcagttaaat cacatactga attagctgaa actgctaatg 1080
gagtgagaat gttagccaaa aatttatccc gagcaaccat tcaattagac gttagagcta 1140
ttatgattat cactaaagct agagataatg gacttattta tttaacaaaa gaagttgttg 1200
aatggatttt ggatcaacat cctcatataa caatttatgc tgatgagaaa ttagcaaagt 1260
cgaaaagatt caatccggaa agtattattg ccaattatcc aaatggttgt aagaaattaa 1320
aatattggaa taaaaaatta actacgaaaa atccagaaat tttcgattta gtacttacat 1380
taggtggtga tggtactgta ttatttgctt caaacttatt tcaaaaaatt gttccaccta 1440
tactttcatt ttcattgggc tcattaggtt ttttaaccaa ttttgaattc agtgcattta 1500
gaacagtatt gagcaaatgt tttgattctg gagttaaagc aaatttgcgt atgcgattca 1560
cttgtcgagt acacactgat gaagggaagt tgatttgtga acaacaagtg ttgaatgaat 1620
tggtagttga tagaggacct agcccatatg ttactcattt ggaattatac ggcgatggat 1680
cattgttaac ggttgcccaa gctgatgggt tgattattgc aactccaact ggttcgactg 1740
cttattcatt atctgctggt gggtctttag ttcaccctgg tgtgagtgcc attagtgtta 1800
ctccaatttg tcctcacacc ttatcgttca gacctatact attacctgat gggatgtttt 1860
tgaaggttaa agtcccactg agcagtagag ccactgcgtg gtgttcattc gatggtaaag 1920
tgcgtactga attgaagaaa ggttattatg tcactattca agcttcacca ttccccttac 1980
ctacagtaat gtcttccaaa acagaatata ttgattctgt cagtagaaat ttacattgga 2040
acatcagaga gcaacaaaaa ccatttagtt catatttgaa accagaaacg cgacaaagta 2100
atgaggaacc tgaaataact gaagattttg atattaatta tactgacaat gaacgtgatt 2220
cttctagttc cactcctagt gaagaaagca acgaagaatg tgctaatacc acgacataa 2279
```

```
<210> 394
```

<400> 394

```
Met Ser His Lys Thr Gln Ser Gln Leu Ser Ser Gln Met Lys Asn Leu

1 5 10 15
```

<sup>&</sup>lt;211> 592

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Candida albicans

9:					EP99870141.1											
	Asn	Thr	Pro	Pro 20	Ile	Asp	Phe	Asn	Ser 25	Thr	Ser	Ser	Asn	Asn 30	Thr	Met
	Pro	Ser	Glu 35	Pro	Asn	Ser	Gln	Pro 40	Gln	Gln	Gln	Gln	Ser 45	Gln	Pro	Glu
	Ala	Lys 50	Thr	Glu	Pro	Gln	Thr 55	Ile	Arg	Pro	Ala	Thr 60	Phe	Thr	Thr	Ser
	Gly 65	Asn	Ser	Ser	Ser	Ser 70	Ser	Ile	Ser	Thr	Leu 75	Ser	Ala	Asp	Ile	Ile 80
	Gln	Pro	Leu	His	Gln 85	Leu	Ser	Ile	Asn	Asn 90	Asn	Asn	Ser	Thr	Val 95	Thr
	Gln	Pro	Ala	Pro 100	Gln	Ser	Ser	Ser	Phe 105	Gln	Arg	Arg	Asn	Asn 110	Pro	Gln
	Arg	Phe	Asn 115	Arg	Așn	Gln	Leu	Asn 120	Val	Tyr	Thr	Asp	Phe 125	Asn	Ser	Thr
	Thr	Ser 130	Ser	Ala	Ser	Ser	Ile 135	Ser	Ser	Ser	Pro	Lys 140	Asp	Phe	Phe	Thr
	Arg 145	Glu	Pro	Pro	Arg	Ile 150	His	Ser	Lys	Leu	Ile 155	Cys	Glu	Glu	Ile	Ala 160
	Ser	Ala	Asn	Asn	Arg 165	Ala	Ala	Lys	Glu	Val 170	Leu	Ser	Arg	Leu	Ser 175	Thr
	Asp	Glu	Leu	Arg 180	Ser	Val	Lys	Ser	His 185	Thr	Glu	Leu	Ala	Glu 190	Thr	Ala
	Asn	Glv	Val	Arg	Met	Leu	Ala	Lvs	Asn	Leu	Ser	Ara	Ala	Thr	Ile	Gln

Asn Gly Val Arg Met Leu Ala Lys Asn Leu Ser Arg Ala Thr Ile Gln 

Leu Asp Val Arg Ala Ile Met Ile Ile Thr Lys Ala Arg Asp Asn Gly 

Leu Ile Tyr Leu Thr Lys Glu Val Val Glu Trp Ile Leu Asp Gln His 

Pro His Ile Thr Ile Tyr Ala Asp Glu Lys Leu Ala Lys Ser Lys Arg 

Phe Asn Pro Glu Ser Ile Ile Ala Asn Tyr Pro Asn Gly Cys Lys 

Leu Lys Tyr Trp Asn Lys Lys Leu Thr Thr Lys Asn Pro Glu Ile Phe Asp Leu Val Leu Thr Leu Gly Gly Asp Gly Thr Val Leu Phe Ala Ser Asn Leu Phe Gln Lys Ile Val Pro Pro Ile Leu Ser Phe Ser Leu Gly Ser Leu Gly Phe Leu Thr Asn Phe Glu Phe Ser Ala Phe Arg Thr Val Leu Ser Lys Cys Phe Asp Ser Gly Val Lys Ala Asn Leu Arg Met Arg Phe Thr Cys Arg Val His Thr Asp Glu Gly Lys Leu Ile Cys Glu Gln Gln Val Leu Asn Glu Leu Val Val Asp Arg Gly Pro Ser Pro Tyr Val Thr His Leu Glu Leu Tyr Gly Asp Gly Ser Leu Leu Thr Val Ala Gln Ala Asp Gly Leu Ile Ile Ala Thr Pro Thr Gly Ser Thr Ala Tyr Ser Leu Ser Ala Gly Gly Ser Leu Val His Pro Gly Val Ser Ala Ile Ser Val Thr Pro Ile Cys Pro His Thr Leu Ser Phe Arg Pro Ile Leu Leu Pro Asp Gly Met Phe Leu Lys Val Lys Val Pro Ser Ser Ser Arg Ala Thr Ala Trp Cys Ser Phe Asp Gly Lys Val Arg Thr Glu Leu Lys Lys Gly Tyr Tyr Val Thr Ile Gln Ala Ser Pro Phe Pro Leu Pro Thr Val Met Ser Ser Lys Thr Glu Tyr Ile Asp Ser Val Ser Arg Asn Leu His

Trp Asn Ile Arg Glu Gln Gln Lys Pro Phe Ser Ser Tyr Leu Lys Pro 515 520 525

```
<210> 395
<211> 1042
<212> DNA
<213> Candida albicans
```

<400> 395 atagoggoog ogcatataat agagaatatg toattacoag ottoatttga ottaactooa 60 gaagatgcta aattgttatt agctgccaac gtccatttgg gtgctaagaa cgttcaagtt 120 cacaacaaac catatgttta caaaaccaga ccagatggta tgaacatcat caacattggt 180 aaaacttggg aaaaaattgt tttggctgcc agaattattg ctgctgttcc aaacgcttct 240 gatgttgctg tttgttcttc aagaactttc ggtcaaagag ctgttttgaa atttgctgct 300 cacactggtg ctactgccat tgctggtaga ttcactccag gtaactttac caattatatc 360 actogttcat tcaaagaacc aagattagtt gttgttactg acccaagaac cgatgctcaa 420 gccatcaaag aatcatctta tgttaacatt ccagttattg ccttgactga catgcagtct 480 ccatctgaat acgttgatgt tgccattcca tgtaacaaca aaggtaaaca ctgtattggt 540 ttaatctggt ggttgcttgc tagagaagtc ttgagattaa gaggtattat cccagacaga 600 actaccgaat ggtcagttat gccagatttg tacttctaca gagacccaga agaaattgaa 660 caaaatgccg tcgaagaagc taaaactgaa ggagttgaag gagctccagt tgctgaagct 720 gaaaccgaat ggactggtga aactgaagat gttgattggg ctgattctgg tgctaccccc 780 agctgctgaa gatgctgctg cttctatctg gtaaacactg aaatctacca ataagaagta 840 gaagtagaag tagaagaaga aacaataaca acaataacaa ccaaaataaa aaaaaggttt 900 aatgatgtat attatcgata aggagaaaga agagattttc ttttttaata atgaggatgc 960

```
<210> 396
<211> 253
<212> PRT
<213> Candida albicans
```

ataagtaaaa aaaaaaaaa aa

	I> 35													_	_
Met	Ser	Leu	Pro	Ala	Ser	Phe	Asp	Leu	Thr	Pro	Glu	Ąsp	Ala	Lys	Leu
1				5					10					15	

Leu Leu Ala Ala Asn Val His Leu Gly Ala Lys Asn Val Gln Val His
20 25 30

Asn Lys Pro Tyr Val Tyr Lys Thr Arg Pro Asp Gly Met Asn Ile Ile 35 40 45

Asn Ile Gly Lys Thr Trp Glu Lys Ile Val Leu Ala Ala Arg Ile Ile
50 55 60

Ala Ala Val Pro Asn Ala Ser Asp Val Ala Val Cys Ser Ser Arg Thr 65 70 75 80

Phe Gly Gln Arg Ala Val Leu Lys Phe Ala Ala His Thr Gly Ala Thr 85 90 95

Ala Ile Ala Gly Arg Phe Thr Pro Gly Asn Phe Thr Asn Tyr Ile Thr 100 105 110

Arg Ser Phe Lys Glu Pro Arg Leu Val Val Val Thr Asp Pro Arg Thr 115 120 125

Asp Ala Gln Ala Ile Lys Glu Ser Ser Tyr Val Asn Ile Pro Val Ile 130 135 140

Pro Cys Asn Asn Lys Gly Lys His Cys Ile Gly Leu Ile Trp Trp Leu 165 170 175

Leu Ala Arg Glu Val Leu Arg Leu Arg Gly Ile Ile Pro Asp Arg Thr 180 185 190

Thr Glu Trp Ser Val Met Pro Asp Leu Tyr Phe Tyr Arg Asp Pro Glu 195 200 205

Glu Ile Glu Gln Asn Ala Val Glu Glu Ala Lys Thr Glu Gly Val Glu 210 215 220

Gly Ala Pro Val Ala Glu Ala Glu Thr Glu Trp Thr Gly Glu Thr Glu 225 230 235 240

Asp Val Asp Trp Ala Asp Ser Gly Ala Thr Pro Ser Cys 245 250

```
<210> 397
<211> 1335
<212> DNA
<213> Candida albicans
<400> 397
teggggetae aaatttegte catgaaaatt gggttegeea ceaataettt gtatgegatt 60
atgcatgctc cacgaggtga aaataccgag gcaatggcgt tggttgtgcc atggactaat 120
tctgacaacg agtacaatga aggtgctatg agtttggcgg tggctttggc acggtacttt 180
acaaagatgt cgatctggtc gaaaaacatt atttttgtat ttcctgagac gggccacaga 240
ccgttgaggt cgtgggttga ggcataccat acggtgttgg acgatactgc ggggtcgatt 300
gaggcggcga ttattatgga gtacggcaag aacggtgatt attttgagta ttacgatatg 360
ttctacgaag ggttgaatgg gcagttgccg aatttggact tgttgaatac ggccaatgta 420
atgacgtatc atgaacagat cccctgtgcc atgcaaggga tgtcggatag ggttatcaat 480
tatagcaccc ggttgcagac tttgtttagg ggtatcctca aattgacgct tgtcgggttg 540
actgatgaag ttcatgggtg tgaagcattt tcggggtggc agatccaggc atttacgatc 600
aaggtaaggg ggactgaagg gaaagatgtt acgcagtttg gccggattgt cgattctacg 660
tttaggtcgg ttaacaattt gcttgaaaag tttcaccaat cgtttttctt ttacttgatg 720
ttgtcgccaa aacactttgt gtctattggg acgtacttgc cgtcggcgat tttgttggca 780
gtatcgtatg cgttgagctc tgtcagtgcg gtggtggttg ccgggtttga ttttcgaaag 840
ctatattttg tggtggtggt tgaaattgcg tgtgctattt tggcgtttgt gccggtgaac 900
caggtgatgc ttgtagcgat tctggcggtg gtgttgttgc cgcgccaagc catcttttcc 960
aagcaggcgg cgttttcgct aatttctatt gcgttgttgg cagtggcatt acttattacc 1020
gccctcttga ttgtacattt tgcattggcg tttagtattg ggattttagc ccttccattg 1080
acatttgtcc cgacattaat gaagaacaag tctaggctaa cagctttttg tttggcggtg 1140
tegaateegt tttttgtgat tttegttget gggaaagtge ttggceacce egagetattt 1200
gaccggttgg tcactgcctg gtcggacata cagtgttgga catggtttat cgttgttttg 1260
gggtggttcc cagcgtgggt gattatcaca ctaagctact gtggctacaa gccagttaag 1320
gaaaaaagtg aatag
                                                                   1335
<210> 398
<211> 444
<212> PRT
<213> Candida albicans
<400> 398
Ser Gly Leu Gln Ile Ser Ser Met Lys Ile Gly Phe Ala Thr Asn Thr
                  5
                                     10
Leu Tyr Ala Ile Met His Ala Pro Arg Gly Glu Asn Thr Glu Ala Met
             20
                                 25
Ala Leu Val Val Pro Trp Thr Asn Ser Asp Asn Glu Tyr Asn Glu Gly
```

484

40

Ala Met Ser Leu Ala Val Ala Leu Ala Arg Tyr Phe Thr Lys Met Ser 50 55 60

Ile Trp Ser Lys Asn Ile Ile Phe Val Phe Pro Glu Thr Gly His Arg
65 70 :75 80

Pro Leu Arg Ser Trp Val Glu Ala Tyr His Thr Val Leu Asp Asp Thr 85 90 95

Ala Gly Ser Ile Glu Ala Ala Ile Ile Met Glu Tyr Gly Lys Asn Gly

100 105 110

Asp Tyr Phe Glu Tyr Tyr Asp Met Phe Tyr Glu Gly Leu Asn Gly Gln
115 120 125

Leu Pro Asn Leu Asp Leu Leu Asn Thr Ala Asn Val Met Thr Tyr His 130 135 140

Glu Gln Ile Pro Cys Ala Met Gln Gly Met Ser Asp Arg Val Ile Asn 145 150 155 160

Tyr Ser Thr Arg Leu Gln Thr Leu Phe Arg Gly Ile Leu Lys Leu Thr 165 170 175

Leu Val Gly Leu Thr Asp Glu Val His Gly Cys Glu Ala Phe Ser Gly
180 185 190

Trp Gln Ile Gln Ala Phe Thr Ile Lys Val Arg Gly Thr Glu Gly Lys
195 200 205

Asp Val Thr Gln Phe Gly Arg Ile Val Asp Ser Thr Phe Arg Ser Val 210 215 220

Asn Asn Leu Leu Glu Lys Phe His Gln Ser Phe Phe Phe Tyr Leu Met 225 230 235 240

Leu Ser Pro Lys His Phe Val Ser Ile Gly Thr Tyr Leu Pro Ser Ala 245 250 255

Ile Leu Leu Ala Val Ser Tyr Ala Leu Ser Ser Val Ser Ala Val Val 260 265 270

Val Ala Gly Phe Asp Phe Arg Lys Leu Tyr Phe Val Val Val Glu 275 280 285

Ile Ala Cys Ala Ile Leu Ala Phe Val Pro Val Asn Gln Val Met Leu 290 295 300

Val Ala Ile Ser Ala Val Val Leu Leu Pro Arg Gln Ala Ile Phe Ser 305 310 315 320

Lys Gln Ala Ala Phe Ser Leu Ile Ser Ile Ala Leu Leu Ala Val Ala 325 330 335

Leu Leu Ile Thr Ala Leu Leu Ile Val His Phe Ala Leu Ala Phe Ser 340 345 350

Ile Gly Ile Leu Ala Leu Pro Leu Thr Phe Val Pro Thr Leu Met Lys

365

355 360

Asn Lys Ser Arg Leu Thr Ala Phe Cys Leu Ala Val Ser Asn Pro Phe 370 375 380

Phe Val Ile Phe Val Ala Gly Lys Val Leu Gly His Pro Glu Leu Phe 385 390 395 400

Asp Arg Leu Val Thr Ala Trp Ser Asp Ile Gln Cys Trp Thr Trp Phe 405 410 415

Ile Val Val Leu Gly Trp Phe Pro Ala Trp Val Ile Ile Thr Leu Ser 420 425 430

Tyr Cys Gly Tyr Lys Pro Val Lys Glu Lys Ser Glu
435 440

<210> 399

<211> 1190

<212> DNA

<213> Candida albicans

<400> 399

```
gaggtggtat gcaaatctt gttaaaactt taactggtaa gactatcact ttggaagtcg 780 aatcttctga caccatcgat aacgtcaaat ccaagatcca agacaaagaa ggtattccac 840 cagaccaaca aagattgatt ttcgccggta aacaattgga agacggtaga accttgtctg 900 actacaacat ccaaaaagaa tctactttac atttggtttt aagattgaga ggtggtatgc 960 aaatctttgt taaaacttta actggtaaga ctatcacttt ggaagtcgaa tcttctgaca 1020 ccatcgataa cgtcaaatcc aagatccaag acaaagaagg tattccacca gatcaacaaa 1080 gattgatttt tgctggtaaa caattagaag atggcagaac cttgtctgac tacaacatcc 1140 aaaaaagaatc taccttgcac ttggtcttga gattgagagg tggtttctaa 1190
```

<del><211> 229</del>

<212> PRT

<213> Candida albicans

<400> 400

Met Gln Ile Phe Val Lys Thr Leu Thr Gly Lys Thr Ile Thr Leu Glu

1 5 10 15

Val Glu Ser Ser Asp Thr Ile Asp Asn Val Lys Ser Lys Ile Gln Asp 20 25 30

Lys Glu Gly Ile Pro Pro Asp Gln Gln Arg Leu Ile Phe Ala Gly Lys 35 40 45

Gln Leu Glu Asp Gly Arg Thr Leu Ser Asp Tyr Asn Ile Gln Lys Glu 50 55 60

Ser Thr Leu His Leu Val Leu Arg Leu Arg Gly Gly Met Gln Ile Phe 65 70 75 80

Val Lys Thr Leu Thr Gly Lys Thr Ile Thr Leu Glu Val Glu Ser Ser 85 90 95

Asp Thr Ile Asp Asn Val Lys Ser Lys Ile Gln Asp Lys Glu Gly Ile 100 105 110

Pro Pro Asp Gln Gln Arg Leu Ile Phe Ala Gly Lys Gln Leu Glu Asp 115 120 125

Gly Arg Thr Leu Ser Asp Tyr Asn Ile Gln Lys Glu Ser Thr Leu His 130 135 140

Leu Val Leu Arg Leu Arg Gly Gly Met Gln Ile Phe Val Lys Thr Leu 145 150 155 160

Thr Gly Lys Thr Ile Thr Leu Glu Val Glu Ser Ser Asp Thr Ile Asp 165 170 175

```
Asn Val Lys Ser Lys Ile Gln Asp Lys Glu Gly Ile Pro Pro Asp Gln
180 185 190
```

Gln Arg Leu Ile Phe Ala Gly Lys Gln Leu Glu Asp Gly Arg Thr Leu
195 200 205

Ser Asp Tyr Asn Ile Gln Lys Glu Ser Thr Leu His Leu Val Leu Arg 210 215 220

Leu Arg Gly Gly Phe

225

<210> 401 <211> 2390 <212> DNA <213> Candida albicans

<400> 401

tccagaatag taattaggga ctttaatttt attaggagtg gattgcatag aatatgctat 60 ctaatagata catcttatta tattettatt catgggaaag caccagtata tggaaatget 120 gcccaaggta gtgcacacaa caaactagtt tacaaagtaa aattcgaagt tacgtcaaag 180 cttagaatgg ttcttatagt agttgtctac tctatcaaaa aacccataat ttgtctatat 240 aacgtaagga tcactatatg ctgttgagta agaaatatgt tgcgagccgt accgacaatg 300 gattggctga aacttgttct catcaaactt gcaaaacact tggagacgcg cgcgcgtttc 360 aacacacaat aaacaacacg aaaaataagg tagaaaacaa aaaaaaaata aaaggaactt 420 taaacaaqaa qtaatcccca ttaaaacttg atcaacactt ttagggtttc cgatttcccc 480 attttcttga ctaaaataat atgagaatac tatgtgttgc cgaaaaacca tcgatttcaa 540 aagaggtggc aaacattttg ggaggagggc gaaaaaaagt aagaaactca cgagaaaaat 600 tcatcaaaaa ctacgatttc accttcactt tcaactctga agatgggcca tgtcaagtaa 660 ccatgacttc ggtggctgga catatcacag gacttgattt tgggtctgcc ttttcgtggg 720 gaaattgtgt tcccgggcga ctatttgaag cagacatcaa gaccattatc accaagaaat 780 ctatttatga aaatattgca gaagaggcaa gaaacgctga taagttgatg atctggacag 840 attgtgatag agaaggagaa tacattggat ttgaaattat gaatgctgca agaaaataca 900 ataggaacct tgggttaaac aatatttggc gagctaggtt ttcacatctt gaacgaaatc 960 acattattcg agcagcaaaa aatcccgtga atttggatat gagtgcagtt tctgcagttt 1020 cttgtcgtat ggaaatcgat cttcgagtgg gtaccagttt tacacgtttg ttgactgatc 1080 aattgagaca aaaggggata attgaaaaga atgaactagc ttcttatggt acatgtcaat 1140 tcccgacatt ggggtttgtt gttgatcgat acaaacgagt caagagtttt acaccggaac 1200 cattctggta tattgagatt gaaactagga aagagaataa aaagacaatt ttcaattggg 1260 ttcgaggtca ttttttcgac aagatgtatg tggttatgct ttatgatcga tgctgcaaaa 1320 gtggagaatt tggaaccata tcaaaaatag aatcaaaacg gaaaccaaat ttccgtccat 1380 tcccattgac aaccgtggag ttacaaaaag attgtgctag attttttaag atgtctgcta 1440 agacggcgtt ggcagctgct gaaagacttt ataacctagg gtatttgtcg tatcctagaa 1500 ctgaaactga caggtttgcc aaagaaaccg atttcaagag cttactagag gtgcacaaac 1560 aagatccgcg atggggaagc tatacaacaa agcttttgaa cgaaggtttc gaaactcctc 1620

```
gaagcggttc tcatgatgat aaggcgcatc ctccaatcca tcctatcaaa tatgtttctt 1680 tggacaccct aaacaccctc gatgaaaaga aagtgtatga atacgttgtg cgacgcttta 1740 ttgcctgttg ctccaaagat gctgttggta cgcaaaccgt ggtgacttta aaatggggag 1800 atgaattctt caccgcaagt ggattaatgg tgcatgaaaa aaattatttg gaagtgtata 1860 cttacaaaaa atgggaaagc tctaaacaac taccgaaatt tacagaggga gaacaggtca 1920 agttgtcgag tggaatattg aaagacggta aaacaagtcc acccaatcat atgaccgagc 1980 ccgagctaat tgcattgatg gatgccaacg gtattggaac cgatgctact atcgctgaac 2040 atattaacaa aatagagact aggcactata ttaataaatt gaaaaagggg aaaaatgaat 2100 atattctcc tactcctta ggaatgggc ttatagaagg ccttgaaaaa atggaatttg 2160 aagatgtatc actatcgaaa ccatttttgc ggaagtcgtt ggaacgatca cttgaggaca 2220 tagcaaccgg gtcccggcca aaagtggatg ttttgaatac aacaataggc gtatatgttg 2280 acgettatag tgtttgttct catcagatac ttgttttgtg caatgaatgt aggagaatta 2340 tacttggaaa tagcagtaac aacaacaaca acaataataa taatacgtaa 2390
```

```
<210> 402
```

<211> 629

<212> PRT

<213> Candida albicans

<400> 402

Met Arg Ile Leu Cys Val Ala Glu Lys Pro Ser Ile Ser Lys Glu Val 1 5 10 15

Ala Asn Ile Leu Gly Gly Gly Arg Lys Lys Val Arg Asn Ser Arg Glu 20 25 30

Lys Phe Ile Lys Asn Tyr Asp Phe Thr Phe Thr Phe Asn Ser Glu Asp 35 40 45

Gly Pro Cys Gln Val Thr Met Thr Ser Val Ala Gly His Ile Thr Gly
50 55 60

Leu Asp Phe Gly Ser Ala Phe Ser Trp Gly Asn Cys Val Pro Gly Arg
65 70 75 80

Leu Phe Glu Ala Asp Ile Lys Thr Ile Ile Thr Lys Lys Ser Ile Tyr 85 90 95

Glu Asn Ile Ala Glu Glu Ala Arg Asn Ala Asp Lys Leu Met Ile Trp
100 105 110

Thr Asp Cys Asp Arg Glu Gly Glu Tyr Ile Gly Phe Glu Ile Met Asn 115 120 125

Ala Ala Arg Lys Tyr Asn Arg Asn Leu Gly Leu Asn Asn Ile Trp Arg . 130 135 140

Ala 145	Arg	Phe	Ser	His	Leu 150	Glu	Arg	Asn	His	Ile 155	Ile	Arg	Ala	Ala	Lys 160
Asn	Pro	Val	Asn	Leu 165	Asp	Met	Ser	Ala	Val 170	Ser	Ala	Val	Ser	Cys 175	Arg
Met	Glu	Ile	Asp 180	Leu	Arg	Val	Gly	Thr 185	Ser	Phe	Thr	Arg	Leu 190	Leu	Thr
Asp	Gln	Leu 195	Arg	Gln	Lys	Gly	Ile 200	Ile	Glu	Lys	Asn	Glu 205	Leu	Ala	Ser
Tyr	Gly 210	Thr	Cys	Gln	Phe	Pro 215	Thr	Leu	Gly	Phe	Val 220	Val	Asp	Arg	Tyr
Lys 225	Arg	Val	Lys	Ser	Phe 230	Thr	Pro	Glu	Pro	Phe 235	Trp	Tyr	Ile	Glu	11e 240
Glu	Thr	Arg	Lys	Glu 245	Asn	Lys	Lys	Thr	Ile 250	Phe	Asn	Trp	Val	Arg 255	Gly
His	Phe	Phe	Asp 260	Lys	Met	Tyr	Val	Val 265	Met	Leu	Tyr	Asp	Arg 270	Суз	Cys
Lys	Ser	Gly 275	Glu	Phe	Gly	Thr	Ile 280	Ser	Lys	Ile	Glu	Ser 285	Lys	Arg	Lys
Pro	Asn 290	Phe	Arg	Pro	Phe	Pro 295	Leu	Thr	Thr	Val	Glu 300	Leu	Gln	Lys	Asp
Cys 305	Ala	Arg	Phe	Phe	Lys 310	Met	Ser	Ala	Lys	Thr 315	Ala	Leu	Ala	Ala	Ala 320
Glu	Arg	Leu	Tyr	Asn 325	Leu	Gly	туг	Leu	Ser 330	Tyr	Pro	Arg	Thr	Glu 335	Thr
Asp	Arg	Phe	Ala 340	Lys	Glu	Thr	Asp	Phe 345	Lys	Ser	Leu	Leu	Glu 350	Val	His
Lys	Gln	Asp 355	Pro	Arg	Trp	Gly	Ser 360	Tyr	Thr	Thr	Lys	Leu 365	Leu	Asn	Glu
Gly	Phe 370	Glu	Thr	Pro	Arg	Ser 375	Gly	Ser	His	Asp	Asp 380	Lys	Ala	His	Pro
Pro 385	Ile	His	Pro	Ile	Lys 390	Tyr	Val	Ser	Leu	Asp 395	Thr	Leu	Asn	Thr	Leu 400

SEQL

Pro Lys Phe Thr Glu Gly Glu Gln Val Lys Leu Ser Ser Gly Ile Leu 465 470 475 480

Lys Asp Gly Lys Thr Ser Pro Pro Asn His Met Thr Glu Pro Glu Leu 485 490 495

Ile Ala Leu Met Asp Ala Asn Gly Ile Gly Thr Asp Ala Thr Ile Ala 500 505 510

Glu His Ile Asn Lys Ile Glu Thr Arg His Tyr Ile Asn Lys Leu Lys 515 520 525

Lys Gly Lys Asn Glu Tyr Ile Leu Pro Thr Pro Leu Gly Met Gly Leu 530 535 540

Ile Glu Gly Leu Glu Lys Met Glu Phe Glu Asp Val Ser Leu Ser Lys 545 550 555 560

Pro Phe Leu Arg Lys Ser Leu Glu Arg Ser Leu Glu Asp Ile Ala Thr 565 570 575

Gly Ser Arg Pro Lys Val Asp Val Leu Asn Thr Thr Ile Gly Val Tyr 580 585 590

Val Asp Ala Tyr Ser Val Cys Ser His Gln Ile Leu Val Leu Cys Asn 595 600 605

Glu Cys Arg Arg Ile Ile Leu Gly Asn Ser Ser Asn Asn Asn Asn 610 615 620

Asn Asn Asn Asn Thr 625

<210> 403

<211> 3098 <212> DNA

```
<213> Candida albicans
<400> 403
atatatagaa ttatggctta gtgcccttta ttaactaaat tagaggttac attaatacaa 60 ·
cttaacaaac aaggaaacta acatccacgt atagggcttc ttatattaat atacctaatt 120
caacctgatt atattgcctt atatacagct tgtaaaaaaa cacataagaa agtttacatc 180
tcaagatgaa ttatccccct attgtaaaaa gtacatcgcc taatgataga catcatagag 240
tactggcgac tccttacatg atggtaacga caacaactac aacaaaaaa aaacattatc 300
ttgacggtat aattagtagt gtgcgagagg cacacgataa gatttatcag tttccttttc 360
teggttgaee ttaatetgtt tttgtataga etttattttt tttgtttttg accacaccca 420
ctttttaata tcacaagata tttaactgat tatagaaaac aacaacaata acccaaatac 480
gttaaccact tttattacat atgatagaca atataatcaa taatttgcaa atcatactac 540
agcaaaatga tgataatttt acatcccctc acgacgatgt gatatatcga ccacattctg 600
ctcgtgtagc acgatatcaa gtaataattg catctacatt gggactcact gccctattac 660
tattttctat cttacgatta aaatatccca aaatatatgt ggcaaacttt aatcatttga 720
atttcagtct acattcgacc tcaagaagga atttacctga attgccttca aattcattat 780
ttggttggat tcctacagtt tacaaaatta ctgagcaaga aattttggaa catgctggat 840
tagatgcagt tgtgtttttg gaatttttta aaatgtgcat tcgaataata agcatatgtt 900
tagtatttgc cattattatc atatctccta tcagatacaa gtttacaggg agagtagatg 960
aagattatcc cgacgatgat agtgacaacg atgacgatga tggaagtaat aataatggta 1020
ccacaataat taagcatata gtgtcagctg gaattctggt ggcaagtaaa aataacgatg 1080
gagaacagta tcaacaattt ctttggttat acactatttt cacctatgta tttacatttg 1140
ttactgttta ctttctattc aaacaaacca ataggattat ttctatgcgt cagaagtatc 1200
ttggatcgca aaattcagtc accgatagaa cggtaaaaat atctgggata ccaggatcat 1260
tacgagatga agtggcactc gcacggcaca ttgaccgttt gaatattggt gaagtggatt 1320
ctgtgttgat tgtcaaggag tggcaaaatc taaacaaact attcaaaaga aggagaagaa 1380
tagttcgaaa attagaggaa agttgggttg aatattttga aaaaaatggg ataaccaata 1440
agagegattt gatategttg cateeccaag ttggegaact gtategtttt tecaataggt 1500
atactgatga tgcagaagaa tcaccagact ggggatctca aaattcgaac tctgcacaag 1560
cttcaataat agatcaagac tcagaatctg ttgaaggaga ttcttctgac actttgaatc 1620
gcttgttgaa tgatgaactg agaacaagac caagtcttcg gaaaggttgg ttcggattgt 1680
ttggacctaa agtggattca ataaactact acaccgataa attggaggtc atagataaag 1740
agattaccag ggccagaact agagaatatc ccgccacttc gactgcattt ctcactatga 1800
aaaccgtggc tgaagcacaa atgttggcac aggcagtctt ggatccaaaa gtcaatcatc 1860
ttatcaccaa cttggcccct gctcctcatg atatccgatg ggataatttg tcattaacta 1920
gacaagacag aaatacaaag atccttgctg tcacgatatt tattggtata atgagtttgt 1980
tgttggttta tccagtcaga tttatggcta gttttttaaa cactaaaagc atttctaaaa 2040
tatggccatc attggggaaa gctattgaat cgcataaatg ggccgaaact ttgattactg 2100
gattattgcc aacttattta tttacaattt taaatattgt gattccgttt ttttatgtat 2160
ggatttctga aaagcaaggt tatttatctc atagtgatga agagttgtca tcggtatcca 2220
agaacttttt ctatatattt gtgaacttat ttttggtttt cacaactttt ggtaccgcct 2280
cttttgttga tacgaccaaa attgcatttg atttagcaag atcactcaga gatttgtcaa 2340
tgttctatgt tgacttaata attctacaag gattgggtat attcccattc aaattattat 2400
tggtggggaa cttacttcgc tttttagtga attcattatt taggtgcaag accccaagag 2460
attacttgaa tttgtacaaa ccgccagttt tcaattttgg tctacaatta ccacaaccaa 2520
tattgatatt tattattacg ttggtatact cggtaatgtc ttcgaagata ttaactgcag 2580
```

```
ggttactata ttttattatt ggttatttg tgagcaaata ccaattgett tatgcttgtg 2640 ttcatcctcc acactcaacg ggcaaagttt ggccaataat tttccgaaga atcatattag 2700 ggttatttct ttttcaaatc acaatggttg gcacattagc attacaagat gcaatcacat 2760 gtgctacttt cttggcacca ttgccgtttt tgacacttta cttttggtgg agtttccata 2820 aacaatatat tcccttgtca acattcattg ccttgagagc aattgaaagc aatgaaaata 2880 tcaaccctac tgattggag caaatcattg aaaacaacaa taataaaact cttgacgaaa 2940 gaagggaatt ggacactaaa tacgaatacc ccaatctagt taatgattta gacgggccaa 3000 tgattgcatt ggatggtgaa gatgattga tagttaatcg agatggtaca actgacgga 3060 aaccacctca atattcagt tcagaatggg actattaa 3098
```

```
<210> 404
```

<211> 865

<212> PRT

<213> Candida albicans

<400> 404

Met Ile Asp Asn Ile Ile Asn Asn Leu Gln Ile Ile Leu Gln Gln Asn 1 5 10 15

Asp Asp Asn Phe Thr Ser Pro His Asp Asp Val Ile Tyr Arg Pro His 20 25 30

Ser Ala Arg Val Ala Arg Tyr Gln Val Ile Ile Ala Ser Thr Leu Gly
35 40 45

Leu Thr Ala Leu Leu Leu Phe Ser Ile Leu Arg Leu Lys Tyr Pro Lys 50 55 60

Ile Tyr Val Ala Asn Phe Asn His Leu Asn Phe Ser Leu His Ser Thr
65 70 75 80

Ser Arg Arg Asn Leu Pro Glu Leu Pro Ser Asn Ser Leu Phe Gly Trp 85 90 95

Ile Pro Thr Val Tyr Lys Ile Thr Glu Glu Ile Leu Glu His Ala 100 105 110

Gly Leu Asp Ala Val Val Phe Leu Glu Phe Phe Lys Met Cys Ile Arg 115 120 125

Ile Ile Ser Ile Cys Leu Val Phe Ala Ile Ile Ile Ile Ser Pro Ile 130 135 140

Arg Tyr Lys Phe Thr Gly Arg Val Asp Glu Asp Tyr Pro Asp Asp Asp 145 150 155 160

Ser Asp Asp Asp Asp Asp Gly Ser Asn Asn Asn Gly Thr Thr Ile

170

175

Ile Lys His Ile Val Ser Ala Gly Ile Ser Val Ala Ser Lys Asn Asn 180 185 190 .

Asp Gly Glu Gln Tyr Gln Gln Phe Leu Trp Leu Tyr Thr Ile Phe Thr 195 200 205

Tyr Val Phe Thr Phe Val Thr Val Tyr Phe Leu Phe Lys Gln Thr Asn 210 215 220

Arg Ile Ile Ser Met Arg Gln Lys Tyr Leu Gly Ser Gln Asn Ser Val
225 230 235 240

Thr Asp Arg Thr Val Lys Ile Ser Gly Ile Pro Gly Ser Leu Arg Asp 245 250 255

Glu Val Ala Leu Ala Arg His Ile Asp Arg Leu Asn Ile Gly Glu Val 260 265 270

Asp Ser Val Leu Ile Val Lys Glu Trp Gln Asn Leu Asn Lys Leu Phe 275 280 285

Lys Arg Arg Arg Ile Val Arg Lys Leu Glu Glu Ser Trp Val Glu 290 295 300

Tyr Phe Glu Lys Asn Gly Ile Thr Asn Lys Ser Asp Leu Ile Ser Leu 305 310 315 320

His Pro Gln Val Gly Glu Ser Tyr Arg Phe Ser Asn Arg Tyr Thr Asp 325 330 335

Asp Ala Glu Glu Ser Pro Asp Trp Gly Ser Gln Asn Ser Asn Ser Ala 340 345 350

Gln Ala Ser Ile Ile Asp Gln Asp Ser Glu Ser Val Glu Gly Asp Ser 355 360 365

Ser Asp Thr Leu Asn Arg Leu Leu Asn Asp Glu Ser Arg Thr Arg Pro

Ser Leu Arg Lys Gly Trp Phe Gly Leu Phe Gly Pro Lys Val Asp Ser 385 390 395 400

Ile Asn Tyr Tyr Thr Asp Lys Leu Glu Val Ile Asp Lys Glu Ile Thr
405 410 415

Arg Ala Arg Thr Arg Glu Tyr Pro Ala Thr Ser Thr Ala Phe Leu Thr

425



Met Lys Thr Val Ala Glu Ala Gln Met Leu Ala Gln Ala Val Leu Asp 435 440 445

Pro Lys Val Asn His Leu Ile Thr Asn Leu Ala Pro Ala Pro His Asp 450 455 460

Ile Arg Trp Asp Asn Leu Ser Leu Thr Arg Gln Asp Arg Asn Thr Lys 465 470 475 480

Ile Leu Ala Val Thr Ile Phe Ile Gly Ile Met Ser Leu Leu Val
485 490 495

400 400

Tyr Pro Val Arg Phe Met Ala Ser Phe Leu Asn Thr Lys Ser Ile Ser 500 505 510

Lys Ile Trp Pro Ser Leu Gly Lys Ala Ile Glu Ser His Lys Trp Ala 515 520 525

Glu Thr Leu Ile Thr Gly Leu Leu Pro Thr Tyr Leu Phe Thr Ile Leu 530 535 540

Asn Ile Val Ile Pro Phe Phe Tyr Val Trp Ile Ser Glu Lys Gln Gly 545 550 555 560

Tyr Leu Ser His Ser Asp Glu Glu Leu Ser Ser Val Ser Lys Asn Phe 565 570 575

Phe Tyr Ile Phe Val Asn Leu Phe Leu Val Phe Thr Thr Phe Gly Thr 580 585 590

Ala Ser Phe Val Asp Thr Thr Lys Ile Ala Phe Asp Leu Ala Arg Ser 595 600 605

Leu Arg Asp Leu Ser Met Phe Tyr Val Asp Leu Ile Ile Leu Gln Gly 610 620

Leu Gly Ile Phe Pro Phe Lys Leu Leu Leu Val Gly Asn Leu Leu Arg 625 630 635 640

Phe Leu Val Asn Ser Leu Phe Arg Cys Lys Thr Pro Arg Asp Tyr Leu 645 650 655

Asn Leu Tyr Lys Pro Pro Val Phe Asn Phe Gly Leu Gln Leu Pro Gln 660 665 670

Pro Ile Leu Ile Phe Ile Ile Thr Leu Val Tyr Ser Val Met Ser Ser

680

685

Lys Ile Leu Thr Ala Gly Leu Leu Tyr Phe Ile Ile Gly Tyr Phe Val 690 695 700

Ser Lys Tyr Gln Leu Leu Tyr Ala Cys Val His Pro Pro His Ser Thr 705 710 715 720

Gly Lys Val Trp Pro Ile Ile Phe Arg Arg Ile Ile Leu Gly Leu Phe 725 730 735

Leu Phe Gln Ile Thr Met Val Gly Thr Leu Ala Leu Gln Asp Ala Ile
740 745 750

Thr Cys Ala Thr Phe Leu Ala Pro Leu Pro Phe Leu Thr Leu Tyr Phe
755 760 765

Trp Trp Ser Phe His Lys Gln Tyr Ile Pro Leu Ser Thr Phe Ile Ala 770 775 780

Leu Arg Ala Ile Glu Ser Asn Glu Asn Ile Asn Pro Thr Asp Leu Glu 785 790 795 800

Gln Ile Ile Glu Asn Asn Asn Asn Lys Thr Leu Asp Glu Arg Arg Glu 805 810 815

Leu Asn Thr Lys Tyr Glu Tyr Pro Asn Leu Val Asn Asp Leu Asp Gly 820 825 830

Pro Met Ile Ala Leu Asp Gly Glu Asp Val Leu Ile Val Asn Arg Asp 835 840 845

Gly Thr Thr Val Arg Lys Pro Pro Gln Tyr Phe Ser Ser Glu Trp Asp 850 855 860

Tyr 865

<210> 405

<211> 1559

<212> DNA

<213> Candida albicans

<400> 405

ttccagaatt agtagatatt tctgatatca agttggagat agggatatcg aactgggatt 60 ctatattttg ttttatcttg gagaacctgt gttgttgttg gcgttctatc tctttatata 120

```
t gaaacatttg aaggaaattc tttcttaada gcatctagtg 180
ttttctctat taattc
acacatgate tetaatetee agtettttga ttaaatatte ttttagaata teaggtgaag 240
aagtgttggt agtcatagct agtagatact gtgtggtgtc tgggataata attataatgt 300
aaacaaaaca aagtcgtgtc agtgtatatt tttcttttgt caattccatc tttttttt 360
tctcttgaga aatgtataac agaggatcca tccatttgct tgacagagaa atacagaaca 420
ctaaacaaac atttttcat tcctttcttg ttttgttctg ttatacccca aaagtttgaa 480
taagtettea gatatetage atggeaactt eteaagaatt gacageagae atacaagete 540
ttgcaactag tttccccaag cgattagcta atgatagtga caattcatta cttattaatg 600
ttgcaccaac tggtcgacaa gccaaaagac atattcaaca gattaattac tccgaagagt 660
ttggagatga ccttgatttt gatgaattcc catcttcgac acccggtact agaagcttaa 720
atgaaaataa agctcaaata gaagcacaaa gatattctct tgcgaaaaac acaccaacgc 780
ccaagagaat cttagaaaaa ccggtgttat ctgagttggt tgaaaaacca gtggtgctta 840
ttcctatcaa aataatgatt gagaatttga acacaaacca aaagttgatt gattccttta 900
tgtggaactt gaatgaaagc ttgattacac caactgagtt tgcggaaatt gtttgcagtg 960
atttagattt accattcagt atggctgcac aaatagcaga ctccattaat caacagattg 1020
aagagtatto ctatgcatct aatttacaac taccaaataa gggcccttac aatgttacca 1080
ttgatttatc agtaaactta aataaacaat tataccaaga tagatttgaa tgggatatga 1140
atcaaaatga agttacacca gaaatttttg ctgaaatagt tgttgctgat ttggggttat 1200
cgttagaatt taagaatgcc atatcacatg cattgcacga aataattatc agagtgaaaa 1260
aagaagtaat agatggtact tttgacaatg aaatgcacaa cttgcatcta gtaaaaggta 1320
taatgtttga acaaggaatt agaattttca ctgaaaacag tgttcaaaat ggaaatgacc 1380
gttgggaacc tttggtcgaa gtattgactt ctagtgaaat cgaaagaaga gaaaatgaaa 1440
gggttagaaa cttgagaaga ttaaagagag agaatatgag aagagattac gatgatcata 1500
gtaggagaag gcaagcagga aaaagaaggt atgatgagtt agaaggagcc tgggtatag 1559
```

<210> 406

<211> 352

<212> PRT

<213> Candida albicans

<400> 406

Met Ala Thr Ser Gln Glu Leu Thr Ala Asp Ile Gln Ala Leu Ala Thr 1 5 10 15

Ser Phe Pro Lys Arg Leu Ala Asn Asp Ser Asp Asn Ser Leu Leu Ile 20 25 30

Asn Val Ala Pro Thr Gly Arg Gln Ala Lys Arg His Ile Gln Gln Ile 35 40 45

Asn Tyr Ser Glu Glu Phe Gly Asp Asp Leu Asp Phe Asp Glu Phe Pro 50 55 60

Ser Ser Thr Pro Gly Thr Arg Ser Leu Asn Glu Asn Lys Ala Gln Ile
65 70 75 80

Glu Ala Gln Arg Tyr Ser Leu Ala Lys Asn Thr Pro Thr Pro Lys Arg

90

95

Ile Leu Glu Lys Pro Val Leu Ser Glu Leu Val Glu Lys Pro Val Val
100 105 110

Leu Ile Pro Ile Lys Ile Met Ile Glu Asn Leu Asn Thr Asn Gln Lys
115 120 125

Leu Ile Asp Ser Phe Met Trp Asn Leu Asn Glu Ser Leu Ile Thr Pro 130 135 140

Thr Glu Phe Ala Glu Ile Val Cys Ser Asp Leu Asp Leu Pro Phe Ser

145 150 155 160

Met Ala Ala Gln Ile Ala Asp Ser Ile Asn Gln Gln Ile Glu Glu Tyr 165 170 175

Ser Tyr Ala Ser Asn Leu Gln Leu Pro Asn Lys Gly Pro Tyr Asn Val 180 185 190

Thr Ile Asp Leu Ser Val Asn Leu Asn Lys Gln Leu Tyr Gln Asp Arg 195 200 205

Phe Glu Trp Asp Met Asn Gln Asn Glu Val Thr Pro Glu Ile Phe Ala 210 215 220

Glu Ile Val Val Ala Asp Leu Gly Leu Ser Leu Glu Phe Lys Asn Ala 225 230 235 240

Ile Ser His Ala Leu His Glu Ile Ile Ile Arg Val Lys Lys Glu Val 245 250 255

Ile Asp Gly Thr Phe Asp Asn Glu Met His Asn Leu His Leu Val Lys
260 265 270

Gly Ile Met Phe Glu Gln Gly Ile Arg Ile Phe Thr Glu Asn Ser Val 275 280 285

Gln Asn Gly Asn Asp Arg Trp Glu Pro Leu Val Glu Val Leu Thr Ser

Ser Glu Ile Glu Arg Arg Glu Asn Glu Arg Val Arg Asn Leu Arg Arg 305 310 315 320

Leu Lys Arg Glu Asn Met Arg Arg Asp Tyr Asp Asp His Ser Arg Arg 325 330 335

Arg Gln Ala Gly Lys Arg Arg Tyr Asp Glu Leu Glu Gly Ala Trp Val

345



```
<210> 407
<211> 737
<212> DNA
<213> Candida albicans
```

<400> 407
aagaaaagta tagtcaaatt gttatacaag ctaaggagcc ataaactttc tttggacatg 60
ctaatatcgc tcctgaccat tgtttactat attcaacaaa acgaattcaa caaggcaaac 120
gaaagctaca tgaaactaag tatcggcaac gtctgttggc caattggggt ggtcaatgta 180
ggtattcatg caagaagtgc agcactgaaa ataaccggtg cctctaacgt cagtaacata 240
atgctaagcg agtctacacg aagatggatt ataagtataa agaggttgat tagttttaaa 300
gagagagtat ataataatgc acgtgattag tttagtaatt ttttgcgagt tagggctata 360
gccctaagac attcacacaa ctaacaaaaa ggaagttctc acgcacataa cgtgtaaccc 420
acataaagaa agaaaaaaaa attcctttg aaaaaattca catcacgctt taaccacttc 480
aacctataac aactccaacc atggctagag aaatcaagga tatcaaagaa ttcgtcgaat 540
tggctagaag atcagacatc aaatctgcta ttgtcaaagt taacgccaag gtcaatgcta 600
acggtaaaaa attcaaacaa accaaattca aggtcagagg ttcaagatac caatacactt 660
tagttgtcaa cgatgcttct aaagctaaaa aattacaaca atcttacca ccaaccttaa 720
aaatcaccaa cttgtaa

<210> 408 <211> 78 <212> PRT <213> Candida albicans

<400> 408

Met Ala Arg Glu Ile Lys Asp Ile Lys Glu Phe Val Glu Leu Ala Arg

1 5 10 15

Arg Ser Asp Ile Lys Ser Ala Ile Val Lys Val Asn Ala Lys Val Asn 20 25 30

Ala Asn Gly Lys Lys Phe Lys Gln Thr Lys Phe Lys Val Arg Gly Ser 35 40 45

Arg Tyr Gln Tyr Thr Leu Val Val Asn Asp Ala Ser Lys Ala Lys Lys
50 55 60

Leu Gln Gln Ser Leu Pro Pro Thr Leu Lys Ile Thr Asn Leu 65 70 75

<210> 409

SEQL

```
<211> 1348
 <212> DNA
 <213> Candida albicans
 <400> 409
 ataccatgac aagcaagctc agggtcgcgg gttcgagtcc cgcggggagc taattatacc 60
 ctcatttttg aacccaccaa attttctttc acattataat gaaataagag tattccctgg 120
 cttcttttt ttttttttt ggeaatatag agaagactgt aataagtata gctcactaaa 180
 agtctttttt tttctattcg ttttatattt ttttaagaaa tttgatgttg atttggtaaa 240
 tgccaaattt taaatgtgtg ttagggctat agccctaatg tactgtatat gcagtatcag 300
 aaatactttt gttacgcaca gtttgtctta ccaaatacat tatatatata tattttttt 360
 tttttgagta gaggagctac actagaccac agtgcgaaaa attcatctct ctatacactt 420
 actcaatttg aagatattca aatttttttc aaaaaaaaat tccttttgga tcgatactag 480
 atagcatata atcatcaaaa atggccaaga tcagtcaagg tatgaaatag atattcaatt 540
 atatacattg aaaaggagat agaacatcaa acaacaacca ttaagaatta agtttaatac 660
 agtttcaata aagagggttt ttttctcaga acaaaccatt gactgaagta ctacaccaag 720
 aaggtataat gatttcacga tttacctgaa tataaagaac atccttaata ttgaatttca 780
 atattaaaaa tacaatttgg ggatattgat gaaattatgt ataggagatt ccatttttca 840
 ttccttcctt atcttacttt tcctattaaa acaaagacta agaaacattc agtaaaatac 960
 taacaaaaat tecatttata tagaegttte tteatetegt tetaaageta gaaaagetta 1020
 tttcactgct tcatcagttg aaagaagagt tcttttatct gctccattat ccaaagaatt 1080
 aagacaacaa tacaatgtca aatctttgcc aattagacaa aatgatgaag ttttagttgt 1140
 tagaggttct aaaaaaggtt ctgaaggtaa agttaattct gtttatagat tgaaatttgc 1200
 tattcaagtt gataaattac aaaaagaaaa atcaaatggt gcttctgttc caatcaacat 1260
 tcatccatct aaagttgtca ttactaaatt acatttggac aaagatagaa aagctttgat 1320
                                                               1348
 tcaaagaaaa ggtggtaaag ctgaataa
 <210> 410
 <211> 127
 <212> PRT
 <213> Candida albicans
···<400>- 410
 Met Ala Lys Ile Ser Ile Asp Val Ser Ser Ser Arg Ser Lys Ala Arg
                                   10
 Lys Ala Tyr Phe Thr Ala Ser Ser Val Glu Arg Arg Val Leu Leu Ser
             20
                                25
 Ala Pro Leu Ser Lys Glu Leu Arg Gln Gln Tyr Asn Val Lys Ser Leu
```

40

45



Pro Ile Arg Gln Asn Asp Glu Val Leu Val Val Arg Gly Ser Lys Lys 50 55 60

Gly Ser Glu Gly Lys Val Asn Ser Val Tyr Arg Leu Lys Phe Ala Ile
65 70 75 80

Gln Val Asp Lys Leu Gln Lys Glu Lys Ser Asn Gly Ala Ser Val Pro 85 90 95

Ile Asn Ile His Pro Ser Lys Val Val Ile Thr Lys Leu His Leu Asp 100 105 110

Lys Asp Arg Lys Ala Leu Ile Gln Arg Lys Gly Gly Lys Ala Glu 115 120 125

<210> 411

<211> 1631

<212> DNA

<213> Candida albicans

## <400> 411

ttcatctttc gaaatcttcc ctcttggaaa cgaccaagag ttggattcgg tgcccttgac 60 tagcacaact ttgttcaaat ctttattgaa taatgcaaca ccacgaaccg gtatggtcga 120 tttatatttt ccgaatttag ataatgette etgtgggtea ecceatttee aaataagegg 180 gcatttctct aaaatctttg tactgaacgt cttcatcttc atgctttgca aatcgggatt 240 caactgacga acaaagtcag cataaaacca ttgtgcttcc tcaatctgaa acattattct 300 ttcaatggaa gataaatcct cttcgggaac attcaccaca aatctaacta acaaatcttc 360 caacaccga tctagagttt gattagctaa accattgcgt aattggatcg acatatcttg 420 ttattttcta tgattaatct ttctttttc tttctttttc actgaatatt cgtaattaaa 480 attgattcaa aattgtattg atgtttgtat actggataaa caacttgtgt gagatcattc 540 aatgtatttg ctggcggaga aggaggcaag gacgaacttt ttttccttca tactttttt 600 tttctctctc tcttttcttt cagtcacatt gtctgagtgt taaaggctgg cttttctgtg 660 ctgaaccttg tggactggta aacgcacgac atagagtaat atttggagaa accaaaagaa 720 ttttgaaaaa ccgtggtttg aattcgacca cagattggtt ggacgacaaa atgcaaagtg 780 tattcatccg gacttttgct acctcaagaa tagagttcca gcggtaccag ccgcgttttg 840 ttaataccat taaagaaacg gtcaaatcgg cacaagaaaa gtcgtacagt attactagac 900 cattggggtt gtcaaaacca gttttgttaa atcataaatt gtcagacaca tactcattgt 960 ccaatatata cgaggagttg tttggtcaaa aatccaaaga aagaagacag aaacaactag 1020 aaatatttac teeteetgtt tegtaettte gacaagacaa atétttgtat tteeeggatt 1140 ttatagcgaa aacattggca ggtaatcaga gaagtttgta cgactcatta gacaatagat 1200 taagcatagt caaattgttt tcttctgttg ctggtgagca gtgtacccgt tcgtacttta 1260 aggttgaaaa caaagattac tattcccagg attatgatac ctttgtggag gaatatcccc 1320 atacccagat acttgatgtg aatatgccgc aaagttggat caaggggttt gtgacaaact 1380 tgagcacagg aaatttaaga aagactttga agccagctct gagatacgag aattatttca 1440 tcttgcctgg ccacataatg tcagcggaaa ttagagaaca gttgtactgt gataatcaat 1500 gttccgggta tatttatatt gttgattcga tggggaagat aagatgggcg acaagtgggt 1560 atgcaactcc tgaggatttg aaattgatgt ggaaggttgt gaaaggggtg caaagagaaa 1620 tgaccaagta a 1631

<210> 412

<211> 376

<212> PRT

<213> Candida albicans

<400> 412

Met Phe Val Tyr Trp Ile Asn Asn Leu Cys Glu Ile Ile Gln Cys Ile 1 5 10 15

Cys Trp Arg Arg Arg Gln Gly Arg Thr Phe Phe Pro Ser Tyr Phe
20 25 30

Phe Phe Ser Leu Ser Leu Phe Phe Gln Ser His Cys Ser Ser Val Lys 35 40 45

Gly Trp Leu Phe Cys Ala Glu Pro Cys Gly Ser Val Asn Ala Arg His 50 55 60

Arg Val Ile Phe Gly Glu Thr Lys Arg Ile Leu Lys Asn Arg Gly Leu 65 70 75 80

Asn Ser Thr Thr Asp Trp Leu Asp Asp Lys Met Gln Ser Val Phe Ile 85 90 95

Arg Thr Phe Ala Thr Ser Arg Ile Glu Phe Gln Arg Tyr Gln Pro Arg
100 105 110

Phe Val Asn Thr Ile Lys Glu Thr Val Lys Ser Ala Gln Glu Lys Ser 115 120 125

Tyr Ser Ile Thr Arg Pro Leu Gly Leu Ser Lys Pro Val Leu Leu Asn 130 135 140

His Lys Leu Ser Asp Thr Tyr Ser Leu Ser Asn He Tyr Glu-Glu-Leu 145 150 155 160

Phe Gly Gln Lys Ser Lys Glu Arg Arg Gln Lys Gln Leu Asp Tyr Asp 165 170 175

Leu Lys His Ser Pro Ile Tyr Glu Val Lys Ser Phe Glu Asn Thr Lys
180 185 190

Gly Lys Ile Phe The Pro Pro Val Ser Tyr Phe Arg Gln Asp Lys Ser 195 200 205

Leu Tyr Phe Pro Asp Phe Ile Ala Lys Thr Leu Ala Gly Asn Gln Arg 210 215 220

Ser Leu Tyr Asp Ser Leu Asp Asn Arg Leu Ser Ile Val Lys Leu Phe 225 230 235 240

Ser Ser Val Ala Gly Glu Gln Cys Thr Arg Ser Tyr Phe Lys Val Glu 245 250 255

Asn Lys Asp Tyr Tyr Ser Gln Asp Tyr Asp Thr Phe Val Glu Glu Tyr 260 265 270

Pro His Thr Gln Ile Leu Asp Val Asn Met Pro Gln Ser Trp Ile Lys 275 280 285

Gly Phe Val Thr Asn Leu Ser Thr Gly Asn Leu Arg Lys Thr Leu Lys 290 295 300

Pro Ala Ser Arg Tyr Glu Asn Tyr Phe Ile Leu Pro Gly His Ile Met 305 310 315 320

Ser Ala Glu Ile Arg Glu Gln Leu Tyr Cys Asp Asn Gln Cys Ser Gly 325 330 335

Tyr Ile Tyr Ile Val Asp Ser Met Gly Lys Ile Arg Trp Ala Thr Ser 340 345 350

Gly Tyr Ala Thr Pro Glu Asp Leu Lys Leu Met Trp Lys Val Val Lys 355 360 365

Gly Val Gln Arg Glu Met Thr Lys 370 375

<210> 413

<211> 1271

<212> DNA

<213> Candida albicans

<400> 413

ccagtgcgtt ttgtttgtt ccacatcata cacttcactg aaactaaata agtttgttta 60 cattttgaga cttcaggtac gacccagggt tgcgacaaag tttaggtagt ttgtcgtctg 120 aatgtcgcaa caaaataggg ctgtagccct agtcatgtga tgtgaattaa cataacaaga 180 agaattgctg gtgcgcaaaa agattatgtg tattttatgt gcgttgttat cctgcacact 240



```
aaaattgagc agtgtacaca cacacatatt gggctgtatt tttattcttg tttttctgct 300
gttctctcac tgttaagctc taagtgaatt tgtgtgtgct gtaatagtgt gtgtgttcca 360
agtoccaget etcacagata etcacgeacg eccatactae tgaaaattte etgactttet 420
gtatctaaaa attttttact aggaattttt ttcttttacg tttttcactt gtttcatata 480
atcaccaact caagtacaac atggctgtcg gtaaaaacaa gagattgtcc aaaggaaaga 540
aaggattaaa aaagaaggtc gttgacccat tcaccagaaa agattggttt gacatcaaag 600
ctccaaccac ttttgaaaac agaaatgttg gtaaaacttt gatcaacaga tctaccggtt 660
taaaqaatqc cqctgatqgc ttgaaaggta gagttttcga agtttgtttg gccgacttac 720
aaggttccga agaccactct tacagaaaaa tcaaattgag agttgatgaa gttcaaggta 780
aaaacttgtt gaccaacttc catggtttgg atttcacttc tgacaaatta agatcattgg 840
tcagaaaatg gcaatcatta gttgaagcta atgtcactgt taaaacttcc gacgattacg 900
ttttgagagt ttttgccatt-gctttcacca aaagacaacc aaaccaaatc aagaaaacta 960
cttacgctca atcctctaaa ttgagagaag tcagaaagaa gatgattgaa attatgcaaa 1020
gagaagtttc caactgtact ttagctcaat taacttccaa attgattcca gaagtcattg 1080
gccgtgaaat tgaaaaatcc acccaaacca ttttcccatt acaaaatgtc cacatcagaa 1140
aagtcaaatt gttgaaacaa ccaaaattcg acttgggttc attattggct ttgcacggtg 1200
aaggttcaac cgaagaaaaa ggtaagaaag tttcttctgg tttcaaagat gttgttttag 1260
                                                                  1271
aatctgttta a
```

<210> 414

<211> 256

<212> PRT

<213> Candida albicans

<400> 414

Met Ala Val Gly Lys Asn Lys Arg Leu Ser Lys Gly Lys Lys Gly Leu

1 5 10 15

Lys Lys Lys Val Val Asp Pro Phe Thr Arg Lys Asp Trp Phe Asp Ile 20 25 30

Lys Ala Pro Thr Thr Phe Glu Asn Arg Asn Val Gly Lys Thr Leu Ile 35 40 45

Asn Arg Ser Thr Gly Leu Lys Asn Ala Ala Asp Gly Leu Lys Gly Arg
50 55 60

Val Phe Glu Val Cys Leu Ala Asp Leu Gln Gly Ser Glu Asp His Ser 65 70 75 80

Tyr Arg Lys Ile Lys Leu Arg Val Asp Glu Val Gln Gly Lys Asn Leu 85 90 95

Leu Thr Asn Phe His Gly Leu Asp Phe Thr Ser Asp Lys Leu Arg Ser 100 105 110

Leu Val Arg Lys Trp Gln Ser Leu Val Glu Ala Asn Val Thr Val Lys

02-07-1999 | EP99870141.1 | SEQL

115 120

Thr Ser Asp Asp Tyr Val Leu Arg Val Phe Ala Ile Ala Phe Thr Lys
130 135 140

Arg Gln Pro Asn Gln Ile Lys Lys Thr Thr Tyr Ala Gln Ser Ser Lys 145 150 155 160

Leu Arg Glu Val Arg Lys Lys Met Ile Glu Ile Met Gln Arg Glu Val 165 170 175

Ser Asn Cys Thr Leu Ala Gln Leu Thr Ser Lys Leu Ile Pro Glu Val

180 185 190

Ile Gly Arg Glu Ile Glu Lys Ser Thr Gln Thr Ile Phe Pro Leu Gln
195 200 205

Asn Val His Ile Arg Lys Val Lys Leu Leu Lys Gln Pro Lys Phe Asp 210 215 220

Leu Gly Ser Leu Leu Ala Leu His Gly Glu Gly Ser Thr Glu Glu Lys 225 230 235 240

Gly Lys Lys Val Ser Ser Gly Phe Lys Asp Val Val Leu Glu Ser Val 245 250 255

<210> 415

<211> 1517

<212> DNA

<213> Candida albicans

<400> 415

tgatatggat ttttcaaaac taagaaacaa taccaactac aacaacaaga aacataagtg 60 tttgcttaat tcaatgacct cgagtcatat tcccacgttt taagtatgag tgttttacga 120 agttgtggat cctatttaa ataaaacaat aatagtaata aaaaaaaaa ccttctttgc 180 ttttcgagaa tttgtaacac attgtttctt tcttcccaca gcaaccaaat tttatttat 240 ttttcttt gggacttacc cacagttgct caattatgta taacaagggt agaaactctg 300 tgggattccc tccttaaaaa tatagcaatc cttttcttc acaacgattg ctatatgacc 360 cccccctaa gcattcattg cttttatata tatttaataa tgtatttctc ttgttcagga 420 taattatcac tatttgtgac gtttaatttt tacatttctt cttcttctc ttcctattc 480 aacaattaaag aacatttaat atgtatttcc caatcattgt atggttatat gtatctatca 540 cttttgtggt tgccaattat ggttttgatc aatggacaaa tgatgatta aacaattt 600 taaaagaacg taaagttgca ttcaatgatg ccttggagaa tccaaaatta attagtttgg 660

```
ctaatgaaga agctaagaaa ttagaaaaag gttacaagaa agttactgaa gaattaaata 720 acaatttgaa tcctccagat gattcattaa atgattattt gaattttgat tacttatttg 780 ggaaaagaaa agaaaattat tcaattaaag aatggattt tgaaaggtgg ccagtaacca 840 gtttgcaaac ttttttaact caaaataata tccaatatag tgcaaaaggat accaaagatg 900 atttaatcaa taaggttaaa gatcaatttg attctattc taagaaaaat catgggtcta 960 gtttttatcc tggcaattgg ttatatgaat cttggtcaga aaatgatttg aaagattggt 1020 tgaaatctta tggcattgaa tttaatccta gttcaacaaa ggatcaattg gttgaaaaat 1080 taaaagaatt tagttatcaa gccactcatt caattagaga ttccaaagaa tctttatttg 1140 attcattgga ttatttgat aaaaccattt ttgataaaaa aggtcaaatt gaagatgaat 1200 tttccaaacc ttggtcatat tctcaattac gtgaataggc ttatttacat ggattattg 1260 acactaaacc aggaatttac gttgaagatt tggataagga aaaattagtc aagattgccc 1320 aaagttataa gaaatgtttg ttgagtgaca ttcatacttg gttggcaaac actgaaaaga 1380 agtctcaacc ttggatcaca aagggagaac aaaagtctca gaaaaagaag ggtagtaatt 1440 tgattaatga tacattcttt gttggtatta ataattggtc caaggataaa ttgcgtgaat 1500 tgggcaatct tgactaa
```

<210> 416

<211> 338

<212> PRT

<213> Candida albicans

<400> 416

Met Tyr Phe Pro Ile Ile Val Trp Leu Tyr Val Ser Ile Thr Phe Val 1 5 10 15

Val Ala Asn Tyr Gly Phe Asp Gln Trp Thr Asn Asp Asp Leu Lys Gln
20 25 30

Phe Leu Lys Glu Arg Lys Val Ala Phe Asn Asp Ala Leu Glu Asn Pro 35 40 45

Lys Leu Ile Ser Leu Ala Asn Glu Glu Ala Lys Lys Leu Glu Lys Gly
50 55 60

Tyr Lys Lys Val Thr Glu Glu Leu Asn Asn Leu Asn Pro Pro Asp
65 70 75 80

Asp Ser Leu Asn Asp Tyr Leu Asn Phe Asp Tyr Leu Phe Gly Lys Arg 85 90 95

Lys Glu Asn Tyr Ser Ile Lys Glu Trp Ile Phe Glu Ser Trp Pro Val 100 105 110

Thr Ser Leu Gln Thr Phe Leu Thr Gln Asn Asn Ile Gln Tyr Ser Ala 115 120 125

Lys Asp Thr Lys Asp Asp Leu Ile Asn Lys Val Lys Asp Gln Phe Asp

130 135 140

Ser Ile Ser Lys Lys Asn His Gly Ser Ser Phe Tyr Pro Gly Asn Trp 145 150 155 160

Leu Tyr Glu Ser Trp Ser Glu Asn Asp Leu Lys Asp Trp Leu Lys Ser 165 170 175

Tyr Gly Ile Glu Phe Asn Pro Ser Ser Thr Lys Asp Gln Leu Val Glu 180 185 190

Lys Leu Lys Glu Phe Ser Tyr Gln Ala Thr His Ser Ile Arg Asp Ser

195 200 205

Lys Glu Ser Leu Phe Asp Ser Leu Asp Leu Phe Asp Lys Thr Ile Phe 210 215 220

Asp Lys Lys Gly Gln Ile Glu Asp Glu Phe Phe Gln Thr Trp Ser Tyr 225 230 235 240

Ser Gln Leu Arg Glu Trp Leu Tyr Leu His Gly Phe Ile Asp Thr Lys 245 250 255

Pro Gly Ile Tyr Val Glu Asp Leu Asp Lys Glu Lys Leu Val Lys Ile 260 265 270

Ala Gln Ser Tyr Lys Lys Cys Leu Leu Ser Asp Ile His Thr Trp Leu 275 280 285

Ala Asn Thr Glu Lys Lys Ser Gln Pro Trp Ile Thr Lys Gly Glu Gln 290 295 300

Lys Ser Gln Lys Lys Lys Gly Ser Asn Leu Ile Asn Asp Thr Phe Phe 305 310 315 320

Val Gly Ile Asn Asn Trp Ser Lys Asp Lys Leu Arg Glu Leu Gly Asn 325 330 335

Leu Asp

<210> 417

<211> 2243

<212> DNA

<213> Candida albicans

```
<400> 417
atgagttcat agatgatctg tttcactttc aaataacgtg tcaacaaaaa taaagttaga 60
gcatagattt tgccgccact tgtacagatg gatagaattg aatgcaaatt ctgcataaag 120
attaaagtga aaaacaattt ccgaaaaaag aagaaaatcg aacacattag aaaaagaaac 180
gaacaaaaga aaaaaaattt caaattgtag ttgcatgtat ataaaataat ataaaagata 240
tatcaccagc acaactgatt actttttatt tatatcacct gtcaacaaca aatttccaaa 300
taaatacaac tcagaaaaaa cacttactat cttttcttag tttggtttct ataatcttat 360
taaacattct tgcctttcat ccttgattat catattagat cttatcttta atttgtttga 420
aaaaataata ccaataatct tcccattaga acttacaaca caacaacaaa aaaacccatt 480
ctaaatcact attctccatt atgaaaattt tcagattatt ttcactacta atcgtacaat 540
ttatcataaa taccactgtt gcagtatcac ctgtgtcagc agttttacca aaactgagtt 600
tcagtccatt tgattcacca gaattttgtt cacagatcat aactcccact tgtaatacaa 660
ctttcaccta cattgatgaa ttgaataaag atattcgtcc ctacttgtcg gaattagtca 720
agaccctgta tttccgttat tttaaagtca atttagataa acaatgtcgt ttttggaatg 780
ctcaacattt ttgtgctagt gaaaactgtg ctgttgaaat attggaagat ttcaattgga 840
gtcaagtcac caatgaaagt ttgaaacctt caggattagg taagatttca ttacctgaca 900
aatcatcaat tgataattcc attgaaaccg aagaagttca aacttgtgaa gatttagatt 960
atagtgaaat agatgatgac catcattgtg tttatgtcaa cttggtaaat aacccagaaa 1020
gatttactgg gtatggtggt aatcaaagtt ttgatgtttg gaaggctatt tacctggaaa 1080
attgtttccc taatactaat ccaatgtcag tgacaaatga tgccgacaat ggtggtgaac 1140
aatgtatcga aaagaacttg ttttatcgtg ttgttagtgg tatgcatgct tcaattgcag 1200
tacatttatc aagagaatat ttgaattctg aaactggtga attttatcct aatttaaagg 1260
tgtttatgga aagagtaggt atgcataatg acagattatc taatatttat ttcaattatg 1320
ccttagtgtc ccaagctata gttaaattga gtgaaatttt accattgaga gagttcattc 1380
aactggggta tgatgacatt actccagctc aaaagcaaca tttattggct aataatgatg 1440
tcgaatcagt cgaagtttat gatcgtttgt tgttagatga cattattcct agtttggaag 1500
caaatgttgt gtttaatact tccaacttgt ttgataatag caatttgagg gatgaattta 1560
gatcaagatt tagaaacatt tctgccatta tggattgtgt tggttgtgat agatgcagaa 1620
tgtgggggaa aatccaaacc attggttatg gtaccgctct caagatttta tttgaagatg 1680
acaactatga taatcacaat ttgaaattta gaagaattga aattgttgcc ttgattaata 1740
cttttgatcg tttatctaaa tctattgaaa gtattaatat gtttaaagaa atgtatttgc 1800
aacaccttaa agatattgct gaaggattaa cccaacctgg tgtttacgac aaaatacaaa 1860
acaacaaacc aggtaacgga tttgccttcc catttgttag tccattacct cagaaaaaac 1920
ctgaccaaac caacacccc aaaaatcaac aacaaaaaca acctcaagaa actgacaaaa 1980
aaagacttac attagaagaa attgcccaca caaaacctga agatcgaact tttattgaag 2040
acttcagatt atcctttgat gaagtttggc aagcattaag atttgtttta actagttatc 2100
aaagattccc agccgtattg agtagattca cattggttca attgaatgaa tggtggaata 2160
aattgcttgg taaaccaaca gtttatgatt accaaagttc ttttgatgtt gatgccctac 2220
                                                                  2243
aatacagtca agtccttgga taa
```

```
<210> 418
```

## <400> 418

Met Lys Ile Phe Arg Leu Phe Ser Leu Leu Ile Val Gln Phe Ile Ile

<sup>&</sup>lt;211> 580

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Candida albicans

Asn Thr Thr Val Ala Val Ser Pro Val Ser Ala Val Leu Pro Lys Ser Ser Phe Ser Pro Phe Asp Ser Pro Glu Phe Cys Ser Gln Ile Ile Thr Pro Thr Cys Asn Thr Thr Phe Thr Tyr Ile Asp Glu Leu Asn Lys Asp Ile Arg Pro Tyr Leu Ser Glu Leu Val Lys Thr Ser Tyr Phe Arg Tyr Phe Lys Val Asn Leu Asp Lys Gln Cys Arg Phe Trp Asn Ala Gln His Phe Cys Ala Ser Glu Asn Cys Ala Val Glu Ile Leu Glu Asp Phe Asn Trp Ser Gln Val Thr Asn Glu Ser Leu Lys Pro Ser Gly Leu Gly Lys Ile Ser Leu Pro Asp Lys Ser Ser Ile Asp Asn Ser Ile Glu Thr Glu Glu Val Gln Thr Cys Glu Asp Leu Asp Tyr Ser Glu Ile Asp Asp Asp His His Cys Val Tyr Val Asn Leu Val Asn Asn Pro Glu Arg Phe Thr Gly Tyr Gly Gly Asn Gln Ser Phe Asp Val Trp Lys Ala Ile Tyr Ser 

Glu Asn Cys Phe Pro Asn Thr Asn Pro Met Ser Val Thr Asn Asp Ala

Asp Asn Gly Glu Gln Cys Ile Glu Lys Asn Leu Phe Tyr Arg Val 

Val Ser Gly Met His Ala Ser Ile Ala Val His Leu Ser Arg Glu Tyr 

Leu Asn Ser Glu Thr Gly Glu Phe Tyr Pro Asn Leu Lys Val Phe Met 

Glu Arg Val Gly Met His Asn Asp Arg Leu Ser Asn Ile Tyr Phe Asn

265

270

Tyr Ala Leu Val Ser Gln Ala Ile Val Lys Leu Ser Glu Ile Leu Pro 275 280 285

Leu Arg Glu Phe Ile Gln Ser Gly Tyr Asp Asp Ile Thr Pro Ala Gln 290 295 300

Lys Gln His Leu Leu Ala Asn Asn Asp Val Glu Ser Val Glu Val Tyr 305 310 315 320

Asp Arg Leu Leu Asp Asp Ile Ile Pro Ser Leu Glu Ala Asn Val 325 330 335

Val Phe Asn Thr Ser Asn Leu Phe Asp Asn Ser Asn Leu Arg Asp Glu 340 345 350

Phe Arg Ser Arg Phe Arg Asn Ile Ser Ala Ile Met Asp Cys Val Gly 355 360 365

Cys Asp Arg Cys Arg Met Trp Gly Lys Ile Gln Thr Ile Gly Tyr Gly 370 380

Thr Ala Leu Lys Ile Leu Phe Glu Asp Asp Asn Tyr Asp Asn His Asn 385 390 395 400

Leu Lys Phe Arg Arg Ile Glu Ile Val Ala Leu Ile Asn Thr Phe Asp 405 410 415

Arg Leu Ser Lys Ser Ile Glu Ser Ile Asn Met Phe Lys Glu Met Tyr
420 425 430

Leu Gln His Leu Lys Asp Ile Ala Glu Gly Leu Thr Gln Pro Gly Val
435 440 445

Tyr Asp Lys Ile Gln Asn Asn Lys Pro Gly Asn Gly Phe Ala Phe Pro 450 455 460

Phe Val Ser Pro Leu Pro Gln Lys Lys Pro Asp Gln Thr Asn Thr Pro 465 470 475 480

Lys Asn Gln Gln Lys Gln Pro Gln Glu Thr Asp Lys Lys Arg Leu 485 490 495

Thr Leu Glu Glu Ile Ala His Thr Lys Pro Glu Asp Arg Thr Phe Ile 500 505 510

Glu Asp Phe Arg Leu Ser Phe Asp Glu Val Trp Gln Ala Leu Arg Phe

525

Val Leu Thr Ser Tyr Gln Arg Phe Pro Ala Val Leu Ser Arg Phe Thr 530 535 540

520

Leu Val Gln Leu Asn Glu Trp Trp Asn Lys Leu Leu Gly Lys Pro Thr 545 550 555 560

Val Tyr Asp Tyr Gln Ser Ser Phe Asp Val Asp Ala Leu Gln Tyr Ser 565 570 575

Gln Val Leu Gly

580

<210> 419

<211> 1004

<212> DNA

<213> Candida albicans

<400> 419

tgtgaaaaaa aattgtggtg tggatgttgt tgtcgttgtt gcgttgtcca caacaaaaa 60 acaagtaaca atttcaaagt tgggcttgga gatcgatttt ttttcccgcg tctgtgtggc 120 acgagacaat tgagtcgacc agtacgtttt aattgaatac gagagtcgac gcaattacat 180 caatccaaca ttccacttat tctatatcaa tgtaaagtca ttttttgata atatcgtaat 240 ttacacattt cgtatatctc ggcaataggg gggataaaaa atagtattga ctaattaata 300 tatcttgttt atcaaatcag gagtatagaa ttccacccaa caactagatt ttccgaatgc 360 gaaacgacga ggacgacaca acaacgacta aagaagaaga agaaaaaaaa tataaataaa 420 ttgatcacgc acacattaga aacacaatat tggatcactt ttttcgataa tactaccacc 480 acacagetea tteaceacte atgeceegaa gttetaetge teaaaagegt ttaetaacag 540 agtatcaaca attatcgagg gacccaccac ctgggataat cgcaggacca gtgagtgaag 600 ataatttata caaatgggaa tgtttattag aaggaccatc cgatactcca tatgcaaatg 660 gagtatecce ageagtattg acttteceta aagattacce attateacca cetacattaa 720 agtttgatcc accattgtta catccaaata tttatgctga tggaaccgtt tgtatttcga 780 ttttacatcc tcctggtgaa gatccaaatc aatatgaacg accagaggaa agatggtcac 840 ctgtgcaaag tattgaaaag atcttgttga gtgtcatgtc tatgcttgca gaacctaatc 900 ctgaaagtgg ggctaatatc gatgcttgta aattatggag agataatcgt gctgaatatg 960 accgacaaat tagacaacat gtcaaggagt cattaggatt atga 1004

<210> 420

<211> 167

<212> PRT

<213> Candida albicans

<400> 420

Met Pro Arg Ser Ser Thr Ala Gln Lys Arg Leu Leu Thr Glu Tyr Gln

1 5 10 15

Gln Leu Ser Arg Asp Pro Pro Pro Gly Ile Ile Ala Gly Pro Val Ser 20 25 30

Glu Asp Asn Leu Tyr Lys Trp Glu Cys Leu Leu Glu Gly Pro Ser Asp 35 40 45

Thr Pro Tyr Ala Asn Gly Val Ser Pro Ala Val Leu Thr Phe Pro Lys
50 55 60

Asp Tyr Pro Leu Ser Pro Pro Thr Leu Lys Phe Asp Pro Pro Leu Leu
65 70 75 80

His Pro Asn Ile Tyr Ala Asp Gly Thr Val Cys Ile Ser Ile Leu His 85 90 95

Pro Pro Gly Glu Asp Pro Asn Gln Tyr Glu Arg Pro Glu Glu Arg Trp
100 105 110

Ser Pro Val Gln Ser Ile Glu Lys Ile Leu Leu Ser Val Met Ser Met
115 120 125

Leu Ala Glu Pro Asn Pro Glu Ser Gly Ala Asn Ile Asp Ala Cys Lys 130 135 140

Leu Trp Arg Asp Asn Arg Ala Glu Tyr Asp Arg Gln Ile Arg Gln His 145 150 155 160

Val Lys Glu Ser Leu Gly Leu 165

<210> 421

<211> 1031

<212> DNA

<213> Candida albicans

## <400> 421

ccctagcctg ccacadeac atgatttctc gtattggatt attgaaaaga cctaccgtgt 540 ccactttaaa caactatgtc aaattacaat cgacattagc ccttaaaaga tacacatcaa 600 ccgtaccagc aacttcaaat caagaacaag aaatattggt tgcccaacgt aaaaatagac 660 ctacatcacc tcatttacaa atttatgaac cacaattaac ttggatcatg tcatcattcc 720 atagaacac tggtgttgct atggccggtg ccttttatgc tttaacttgt ggatttgctg 780 ctacttcaat tttaaatatt ccatttgata ctactacttt agtatctgca ttcaccacat 840 taccaacatt tgctcaatat ggtatcaaag ctatttgtgc ttatccattt gtttatcata 900 ttggtaatgg gattagacat ttggtttggg attttggtaa agaattaacc atccctggtg 960 tttatagaac tgggtatgct gttttggctg ctactgctgt cattggaagt tatttagctt 1020 tcttatggta a

<210> 422

02-07-1999

<211> 176

<212> PRT

<213> Candida albicans

<400> 422

Met Ile Ser Arg Ile Gly Leu Leu Lys Arg Pro Thr Val Ser Thr Leu

1 5 10 15

Asn Asn Tyr Val Lys Leu Gln Ser Thr Leu Ala Leu Lys Arg Tyr Thr
20 25 30

Ser Thr Val Pro Ala Thr Ser Asn Gln Glu Gln Glu Ile Leu Val Ala 35 40 45

Gln Arg Lys Asn Arg Pro Thr Ser Pro His Leu Gln Ile Tyr Glu Pro
50 55 60

Gln Leu Thr Trp Ile Met Ser Ser Phe His Arg Ile Thr Gly Val Ala 65 70 75 80

Met Ala Gly Ala Phe Tyr Ala Leu Thr Cys Gly Phe Ala Ala Thr Ser 85 90 95

Ile Leu Asn Ile Pro Phe Asp Thr Thr Thr Leu Val Ser Ala Phe Thr
100 105 110

Thr Leu Pro Thr Phe Ala Gln Tyr Gly Ile Lys Ala Ile Cys Ala Tyr 115 120 125

Pro Phe Val Tyr His Ile Gly Asn Gly Ile Arg His Leu Val Trp Asp 130 135 140

Phe Gly Lys Glu Leu Thr Ile Pro Gly Val Tyr Arg Thr Gly Tyr Ala 145 150 155 160

Val Leu Ala Ala Thr Ala Val Ile Gly Ser Tyr Leu Ala Phe Leu Trp

165 170 175

<210> 423 <211> 1176 <212> DNA

<213> Candida albicans

<400> 423

aaaagcaaga agagaaggac tcgttggcca atttcttcgg caatttcaag aagaaaagag 60 tagctgttta gaactatata tatatgtact cgcgctttaa tgtttatagc aataatgaaa 120 tgttttaata attatttaat caaacttggt tgtaacttat gattatggta gtgatctaag 180 aacacaactt gcaaagcaat ggtagtttct ttgattttgt gtttctatta gattcctgtt 240 tctattagat tcccgctttt tttttttttg cagacattaa acctcagggc tatagcccta 300 atggcaaaac atgcacgtgt atgtttcttg atttttctac actactagta aaaaaatttt 360 ctttccgctc actattcaca catacactct ttttcgcaca attacagtct accaacagga 420 aaagaaaaaa aaaaggaatc tggtaattga aaaattgaag tttggttctt ttaatactat 480 caatcaacta gagtcacagc atgttaattc caaaagaaga cagaaagaag atccaccaat 540 acctcttcca aggtatgtaa atatgaatta taaactggaa cagaatatgg catttcaagg 600 gatgcacgat aagtcaagag ttcatgaaaa agcacagatt ataacagtcg taaagaaaaa 660 tttcactacc aacaacaata agaagatatc aaagagattc agtaatcact acttacaaga 720 aacatataac atcatggaga gtttaatttg aaatacgaat gaatatacaa atgaactata 780 ccctttttat ggccatatca cgtttcaaga aatatttaaa caaaaataaa atgaagaata 840 aaacttggat atactaacac atgtattata gagggtgtcg ttgttgctaa gaaagacttc 900 aaccaaccaa agcacgatga aattgatact agaaacttgt tcgtcatcaa agctttacaa 960 tctttgactt ctaaaggtta cgtcaagact caattctcat ggcaatacta ctactacacc 1020 ttgactgatg aaggtgttga attcttgaga accgaattga acattccaga aggtatcttg 1080 ccattgacca gattgaagaa tgctccagct gaaagaccaa gaccatcaag aggcggtcca 1140 agaagaggtg gttacagagg tagagctaga gactaa 1176

<210> 424

<211> 118

<212> PRT

<213> Candida albicans

<400> 424

Met Leu Ile Pro Lys Glu Asp Arg Lys Lys Ile His Gln Tyr Leu Phe 1 5 10 15

Gln Glu Gly Val Val Val Ala Lys Lys Asp Phe Asn Gln Pro Lys His
20 25 30

SEQL

Asp Glu Ile Asp Arg Asn Leu Phe Val Ile Lys Ala Leu Gln Ser 35 40 45

Leu Thr Ser Lys Gly Tyr Val Lys Thr Gln Phe Ser Trp Gln Tyr Tyr 50 55 60

Tyr Tyr Thr Leu Thr Asp Glu Gly Val Glu Phe Leu Arg Thr Glu Leu 65 70 75 80

Asn Ile Pro Glu Gly Ile Leu Pro Leu Thr Arg Leu Lys Asn Ala Pro 85 90 95

Ala Glu Arg Pro Arg Pro Ser Arg Gly Gly Pro Arg Arg Gly Gly Tyr

100 105 110

Arg Gly Arg Ala Arg Asp 115

<210> 425

<211> 2840

<212> DNA

<213> Candida albicans

## <400> 425

tgttgggagt attctggcac aacaatact tacttttagg ttactaacat tatttttctt 60 cagctaacta attatctagt ttatatctat atccattatt gttgaaatca ctatcgtgag 120 gtaaataaca actacagagt tgtcacagta tccaaaaaaa actttgtacc tatcaataca 180 ttttaagcaa taggtcattt attgctgtaa tcaagtgtta ccagtatctc tagttaatgt 240 tgagtttata cctaaaacat gaactatatc aactttaaat gcccctaatc acgtgataca 300 gcacatggga atttgctgat cttgcttcct tgcacgtaca cggcacatgt acacgacttt 360 tttttctaaa tattttttt tttgttgttg ttgttattaa acaatattat tttcatatat 420 ataattgctt agtttacttg cttcttacac cctttgcata ttttttttt ttcctttttc 480 tgccaacttg atcaaattcg atgctacatc ctaataattc agtagtcgat atgtctagca 540 ctggaaacat gaatgaaaat acagatgctc caccgaaaca gcagacgaaa aagaaaatat 600 caaaacagaa ttcaaccaaa accgattttt ttgctgcacg attggctagt gcggttgacg 660 atatagaaag cagtgatagt gatgaaacgt ttatatatga gaataatgat actgaacttg 720 atgataatgc tagtaatatt aacaataaca acaataacag caccaataac attatcaatt 780 ctcccggaac atcgatagcg ttaggatcgg gccttcgatc gccatccata ctagaggggg 900 aacagcttca atattttcat gacccagtga ggcaacaaca gttcaaactt ccttctacca 960 aggeteegte aattteeaac teeateagea gtteaaataa tatagattea ataettaaac 1020 gacctgtaca tetacgtgag geatcaacgt atteagtgaa tgataatgac caccgaaate 1080 ttgtgctacc aaactcaact gaaaggttta ctgcttcacc ttcgaacaat attggtaatg 1140 aaaatatacc tcaataccag aaaactagta gtgttgccca ctcaatcaat gaaggataca 1200 atgatgatac attttcatat aatgaagttg aggataattt aattgatgaa gattccacgg 1260 acgatgggga tttgacaaaa aataccatta ctaataacaa caatccacca accacgtcga 1320

```
gccaacagca accacaaccg caaccacagc cgcaaccaca gcaaccacaa ctacatactc 1380
tgtctcctct aaatcagata caagcagcga catcagcaac accttctgtg tccactaaaa 1440
acgcgtctaa aagaaattac aaaacctcat ccacttcctc aaaattaaga tcaactacat 1500
caaaactttt cgataaaaaa gggtcacaac caagaagata cagtaccatt cctgatgata 1560
ttgacattga agatttcgat gatgagctta tatactatga caacacagct aggttcccag 1620
cgaacgaatc aacttcatta ctaaatcaaa accaaagaat cccccattat agatcactta 1680
atttgaattt ccctcaggtg aagcgccaaa gcaagcgtta tttgtcaact ggccaacctt 1740
tagagagttc tgatcgtggc tctaacaaag atggtactga taatggaaac aacagtgatc 1800
acaatattaa ttctcctttg actgctaata ataataataa taacgtcaat cacaacgatc 1860
atggtgataa caaaaagagt aataccaaca acaacaacat tgctaataat agagcatttc 1920
catttcctta tcaagatcaa caacatcatt attactacga ctacgatgat tttgaccaag 1980
aatcacaaat caatggacee-aattttgatt tgccagacct ccctataaac agatcagctt 2040
cacggaattt taacaacaac aataacccca aaagatttgg cgacagtcat ttttttctac 2100
caagaaagac agatcagtat agtcaaagaa caagctttct aaagtcatgc atttatacct 2160
ttgtttgtat attaattgtg cttaccatag ggtttgtatt ggggtttgtt ttggccacaa 2220
caaaagattt aactgatgta ggtatcacat ccattgagaa ccccattgtt agtaaagatg 2280
agttagtttt caatgttgtt attgaagcat ttaatccagg gtggttttcc gttgacatca 2340
atgaagtaga gttggattta tttgcaagaa gtggctatct acctgataca gataatctga 2400
aaatatctaa catgggaggg tcacaaaaag tagagacggt taaattagga acgattttga 2460
attttgaatc ggttctcaat ttcaaaggtg ggtttctctc tcgagaaccg acaattcaaa 2520
agggaggaat ccgattatta tatcctggca aaaatgttac tgccgaggct aaattggttg 2580
taaatatggc tgatattaaa atagctgctt ccaatagcat tgctaaagaa agcactacta 2640
gtaatgacac caacgataat gacaactcca agaagtggga aatcatatca agcaatccat 2700
ttgatttgat tataactgga gtattgaagt atgatttacc attttcaaga acctcaagat 2760
cagtcgtggt tagaaaaact gggtatattg atcctacctt attcgtaata ccacaaggag 2820
                                                                  2840
agaataatat ttcaatttag
```

<210> 426

<211> 779

<212> PRT

<213> Candida albicans

<400> 426

Met Leu His Pro Asn Asn Ser Val Val Asp Met Ser Ser Thr Gly Asn
1 5 10 15

Met Asn Glu Asn Thr Asp Ala Pro Pro Lys Gln Gln Thr Lys Lys
20 25 30

Ile Ser Lys Gln Asn Ser Thr Lys Thr Asp Phe Phe Ala Ala Arg Leu
35 40 45

Ala Ser Ala Val Asp Asp Ile Glu Ser Ser Asp Ser Asp Glu Thr Phe
50 55 60

Ile Tyr Glu Asn Asn Asp Thr Glu Leu Asp Asp Asn Ala Ser Asn Ile
65 70 75 80



02-07-1999

Asn Asn Asn Asn Asn Ser Thr Asn Asn Ile Ile Asn Leu Asp Asn 85 90 95

Ala Ser Val Asn Gly Ser Met Ile Ala Ser Ser Asn Ala Met Val Thr 100 105 110

Gly Pro Pro Gly Thr Ser Ile Ala Leu Gly Ser Gly Leu Arg Ser Pro 115 120 125

Ser Ile Leu Glu Gly Glu Gln Leu Gln Tyr Phe His Asp Pro Val Arg 130 135 140

Gln Gln Gln Phe Lys Leu Pro Ser Thr Lys Ala Pro Ser Ile Ser Asn 145 150 155 160

Ser Ile Ser Ser Ser Asn Asn Ile Asp Ser Ile Leu Lys Arg Pro Val 165 170 175

His Leu Arg Glu Ala Ser Thr Tyr Ser Val Asn Asp Asn Asp His Arg 180 185 190

Asn Leu Val Leu Pro Asn Ser Thr Glu Arg Phe Thr Ala Ser Pro Ser 195 200 205

Asn Asn Ile Gly Asn Glu Asn Ile Pro Gln Tyr Gln Lys Thr Ser Ser 210 215 220

Val Ala His Ser Ile Asn Glu Gly Tyr Asn Asp Asp Thr Phe Ser Tyr 225 230 235 240

Asn Glu Val Glu Asp Asn Leu Ile Asp Glu Asp Ser Thr Asp Asp Gly 245 250 255

Asp Leu Thr Lys Asn Thr Ile Thr Asn Asn Asn Asn Pro Pro Thr Thr 260 265 270

Ser Ser Gln Gln Gln Pro Gln Pro Gln Pro Gln Pro Gln Pro Gln Gln 275 280 285

Pro Gln Leu His Thr Ser Ser Pro Leu Asn Gln Ile Gln Ala Ala Thr 290 295 300

Ser Ala Thr Pro Ser Val Ser Thr Lys Asn Ala Ser Lys Arg Asn Tyr 305 310 315 320

Lys Thr Ser Ser Thr Ser Ser Lys Leu Arg Ser Thr Thr Ser Lys Leu 325 330 335

rue	vsħ	пуз	340	GIY	Ser	Gin	PIO	345	nrg	171	Ser	****	350	110	vah
Asp	Ile	Asp 355	Ile	Glu	Asp	Phe	Asp 360	Asp	Glu	Leu	Ile	Tyr 365	Tyr	Asp	Asn
Thr	Ala 370	Arg	Phe	Pro	Ala	Asn 375	Glu	Ser	Thr	Ser	Leu 380	Leu	Asn	Gln	Asn
Gln 385	Arg	Ile			-								Pro		
Lys	Arg	Gln	Ser	Lys 405	Arg	Tyr	Leu	Ser	Thr 410	Gly	Gln	Pro	Leu	Glu 415	Ser
Ser	Asp	Arg	Gly 420	Ser	Asn	Lys	Asp	Gly 425	Thr	Asp	Asn	Gly	Asn 430	Asn	Ser
Asp	His	Asn 435	Ile	Asn	Ser	Pro	Leu 440	Thr	Ala	Asn	Asn	Asn 445	Asn	Asn	Asn
Val	Asn 450	His	Asn	Asp	His	Gly 455	Asp	Asn	Lys	Lys	Ser 460	Asn	Thr	Asn	Asn
Asn 465	Asn	Ile	Ala	Asn	Asn 470	Arg	Ala	Phe	Pro	Phe 475	Pro	Tyr	Gln	Asp	Gln 480
Gln	His	His	Tyr	туг 485	Tyr	Asp	Tyr	Asp	Asp 490	Phe	Asp	Gln	Glu	Ser 495	Gln
Ile	Asn	Gly	Pro 500	Asn	Phe	Asp	Leu	Pro 505	Asp	Leu	Pro	Ile	Asn 510	Arg	Ser
Ala	Ser	Arg 515	Asn	Phe	Asn	Asn	Asn 520	Asn	Asn	Pro	Lys	Arg 525	Phe	Gly	Asp
Ser	His 530	Phe	Phe	Leu	Pro	Arg 535	Lys	Thr	Asp	Gln	Tyr 540	Ser	Gln	Arg	Thr
Ser 545	Phe	Leu	Lys	Ser	Cys 550	Ile	Tyr	Thr	Phe	Val 555	Суз	Ile	Leu	Ile	Val 560
Leu	Thr	Ile	Gly	Phe 565	Val	Leu	Gly	Phe	Val 570	Leu	Ala	Thr	Thr	Lys 575	Asp
Leu	Thr	Asp	Val	Gly	Ile	Thr	Ser	Ile	Glu	Asn	Pro	Ile	Val	Ser	Lys

Ser Val Leu Asn Phe Lys Gly Gly Phe Leu Ser Arg Glu Pro Thr Ile 660 665 670

Gln Lys Gly Gly Ile Arg Leu Leu Tyr Pro Gly Lys Asn Val Thr Ala 675 680 685

Glu Ala Lys Leu Val Val Asn Met Ala Asp Ile Lys Ile Ala Ala Ser 690 695 700

Asn Ser Ile Ala Lys Glu Ser Thr Thr Ser Asn Asp Thr Asn Asp Asn 705 710 715 720

Asp Asn Ser Lys Lys Trp Glu Ile Ile Ser Ser Asn Pro Phe Asp Leu
725 730 735

Ile Ile Thr Gly Val Leu Lys Tyr Asp Leu Pro Phe Ser Arg Thr Ser 740 745 750

Arg Ser Val Val Val Arg Lys Thr Gly Tyr Ile Asp Pro Thr Leu Phe 755 760 765

Val Ile Pro Gln Gly Glu Asn Asn Ile Ser Ile 770 775

645

<210> 427

<211> 1352

<212> DNA

<213> Candida albicans

<400> 427

cccacagacc aataacgttt taccaaccaa aacctctgat atcaatgatt cgaaaagatt 60 cattcaaccc cggttagctg tttgagtatt ggatagcaac actttcaatt agtgcaacac 120 aatcaaatta ccaatacttg tttacctttc atctgattct aattggttca tagcaatata 180

```
gtctcttctg ttgtttgata ttaatataaa taaaacttat ttatcacgtt gtttagtagt 240
ctcgcaaatt tgaaaccatg gatgagtaaa cttgttgtgt tagatgagct caaatatctg 300
gtggaacaat tgtgtagtag ctctttgata aatatccaag aacagtcgtg caagtttcaa 360°
taatagctca tcacaacaat tcgtctacac ttcccacctg atttgttggt ttaaatataa 480
taagacaaac ctcagaagct atgataagaa aacaggctag agaaagaaga gagtatcttt 540
atagaaaggc tttacagctt caggaatctt ccttaacaga aaaaagacaa caattgaaag 600
cagctctagc aagtggaaaa tcattatcaa aggagcttgc cgaagatgaa aaattacaac 660
qtqattttat ttacqatqaa aqtqaacaaa taqaaattga tgacqaatac agtcgqttgt 720
cgggaatate tgatecaaaa gttgttatta ecacateceg tgatecatet gteaagttge 780
tacaattcct gaaagaaatc aagttaatgt ttccaaatag cttgaagttg aatcgaggaa 840
actatataat ctcagatttg gtaagtacct gtaatagagt gcaagtttcc gatatgattt 900
tattgcacga gcatcgtggt gtcccatcaa gtttaactgt aagccacttt cctcatggcc 960
caactgcgat tttcacgtta cataatgtca aactaagaca cgatttgcca aacttgggaa 1020
acgtctcaga gtcctatcct cacttaatat ttgagaattt ccaatccgac ttgggtaagc 1080
gtgtggttaa aatattgcaa catttgtttc ctccaggtgt caagaaagat agctccagag 1140
taataacatt tgtcaataac gatgactaca tatcggtgag acaccatgtt tacgtcaaaa 1200
ctaaggattc agtggagttg agtgagattg gcccacgttt cgaaatgaga ttgtatgaaa 1260
tcagactagg attacctgac aacaaagatg ctgatgtcga gtggcagatg agaagattca 1320
taagaacagc taatagaaag aattacttgt aa
                                                               1352
```

<210> 428

<211> 283

<212> PRT

<213> Candida albicans

<400> 428

Met Ile Arg Lys Gln Ala Arg Glu Arg Arg Glu Tyr Leu Tyr Arg Lys

1 5 10 15

Ala Leu Gln Leu Gln Glu Ser Ser Leu Thr Glu Lys Arg Gln Gln Leu
20 25 30

Lys Ala Ala Leu Ala Ser Gly Lys Ser Leu Ser Lys Glu Leu Ala Glu 35 40 45

Asp Glu Lys Leu Gln Arg Asp Phe Ile Tyr Asp Glu Ser Glu Gln Ile 50 55 60

Glu Ile Asp Asp Glu Tyr Ser Arg Leu Ser Gly Ile Ser Asp Pro Lys
65 70 75 80

Val Val Ile Thr Thr Ser Arg Asp Pro Ser Val Lys Leu Leu Gln Phe
85 90 95

Ser Lys Glu Ile Lys Leu Met Phe Pro Asn Ser Leu Lys Leu Asn Arg 100 105 110

Gly Asn Tyr Ile Ile Ser Asp Leu Val Ser Thr Cys Asn Arg Val Gln
115 120 125

Val Ser Asp Met Ile Leu Leu His Glu His Arg Gly Val Pro Ser Ser 130 135 140

Leu Thr Val Ser His Phe Pro His Gly Pro Thr Ala Ile Phe Thr Leu 145 150 155 160

His Asn Val Lys Leu Arg His Asp Leu Pro Asn Leu Gly Asn Val Ser 165 170 175

Glu Ser Tyr Pro His Leu Ile Phe Glu Asn Phe Gln Ser Asp Leu Gly
180 185 190

Lys Arg Val Val Lys Ile Leu Gln His Leu Phe Pro Pro Gly Val Lys 195 200 205

Lys Asp Ser Ser Arg Val Ile Thr Phe Val Asn Asn Asp Asp Tyr Ile 210 215 220

Ser Val Arg His His Val Tyr Val Lys Thr Lys Asp Ser Val Glu Leu 225 230 235 240

Ser Glu Ile Gly Pro Arg Phe Glu Met Arg Leu Tyr Glu Ile Arg Leu 245 250 255

Gly Leu Pro Asp Asn Lys Asp Ala Asp Val Glu Trp Gln Met Arg Arg 260 265 270

Phe Ile Arg Thr Ala Asn Arg Lys Asn Tyr Leu 275 280

<210> 429

<211> 1061

<212> DNA

<213> Candida albicans

<400> 429

tccttttgtt tttattttg ctgtgttact ccagaaatgt gcataataat gataatagta 60 atttgtgact aatatgagat gatcgtatgt gggtgggtgg ggaggaaggg acccggaatt 120 ctaggaacag gaaaaataaa aacgaataaa caaaaacccc ccaatcggca tgcatcggaa 180 ttctttcagc ccaattactt tatttttgcc cacttctttt ggattagggc aatagcccta 240 aagctcgtgt tttagccctt tatatgcagt ctattttatt tttctctttt ttttttggct 300 gttggtaaac ttttttttt ttcgcaggtg ttgaaaaaaaa aatcatttt acagtttaca 360

```
tttctctaac ctgcaaaaag ctctcgttt tttgtagtga gagttactcg ttcacaatag 420 tatactttac aggggagttc ttttcttttg gaatagtcaa ccaacagcaa atagccaagg 480 atcaagcttc atcattaatc atgtcctcta agatcttatc agaaaaccca actgaattag 540 aattaaaagt tgctcaagct ttcgttgatt tggaatctca agctgattta aaagctgaat 600 tgagaccatt acaattcaaa tctatcaaag aaattgatgt taatggaggt aaaaaagctt 660 tagctgtttt cgttcacca ccaagtttac aagcttacag aaaagttcaa actagattaa 720 ctagagaatt agaaaaaaaa ttcccagata gacatgttgt ctttttagct gaaagaagaa 780 tcttaccaaa accagctaga aaagctagaa aacaacaaaa aagaccaaga tcaagaactt 840 tgactgctgt tcatgataaa attttggaag atttagttt cccaactgaa atcattggta 900 aaagagttag atacttggtt ggtggtaaca aaatccaaaa agtcttgttg gattctaaag 960 attcaactgc tgttgattac aaattggatt ctttccaaca attgtactca aaattgactg 1020 gtaaaacaagt tgtttttgaa atcccaggtg aatctcatta g
```

```
<210> 430
```

<211> 186

<212> PRT

<213> Candida albicans

<400> 430

Met Ser Ser Lys Ile Leu Ser Glu Asn Pro Thr Glu Leu Glu Leu Lys

1 5 10 15

Val Ala Gln Ala Phe Val Asp Leu Glu Ser Gln Ala Asp Leu Lys Ala 20 25 30

Glu Leu Arg Pro Leu Gln Phe Lys Ser Ile Lys Glu Ile Asp Val Asn 35 40 45

Gly Gly Lys Lys Ala Leu Ala Val Phe Val Pro Pro Pro Ser Leu Gln 50 55 60

Ala Tyr Arg Lys Val Gln Thr Arg Leu Thr Arg Glu Leu Glu Lys Lys
65 70 75 80

Phe Pro Asp Arg His Val Val Phe Leu Ala Glu Arg Arg Ile Leu Pro 85 90 95

Lys Pro Ala Arg Lys Ala Arg Lys Gln Gln Lys Arg Pro Arg Ser Arg 100 105 110

Thr Leu Thr Ala Val His Asp Lys Ile Leu Glu Asp Leu Val Phe Pro 115 120 125

Thr Glu Ile Ile Gly Lys Arg Val Arg Tyr Leu Val Gly Gly Asn Lys 130 135 140

Ile Gln Lys Val Leu Leu Asp Ser Lys Asp Ser Thr Ala Val Asp Tyr

150

155

160

Lys Leu Asp Ser Phe Gln Gln Leu Tyr Ser Lys Leu Thr Gly Lys Gln 165 170 175

Val Val Phe Glu Ile Pro Gly Glu Ser His 180 185

<210> 431

<211> 1256

<212> DNA

<213> Candida albicans

<400> 431

tgttttgttt ttgcaatcaa cataatagaa gaccaaacaa ataatttcta ttttttttga 60 etetecegta gttttacact tetageetet ettgtaaata tacacetaat tgacagtace 120 attaggaccc catcttattg ttaaggataa tacttcttct tcttcttctt ctttgtttaa 180 tcaaatttgc aataaataaa aaaaaaaaa aaacaaagcc gcacaagttt tcctaaaatg 240 acttattttg tgtaacgcat tcacgtgatc ataatttttt taaattcaaa aactgaacca 300 aattcctgca tattgaggtt gaaaaaaaaa agaaaaagaa aattttttca atcttgtttg 360 aggagagaga ggtgaaaaat ttttctctct ctctttcttt ctttcattct catataccat 420 aaacttaaac aacttetttt acttttteet ttetttteet tteaaacete tacaacagat 480 ccaattaatt aacaaaaaa atggttaacg ctatcttatc taagaaaaag aaattagtag 540 ctgacggtgt cttctacgct gaattgaacg aattcttcac cagagaatta gctgaacaag 600 gttatgctgg tgttgaagtt agaaaaactc catctaaatt ggaagttatt gttaaagctt 660 ctaacactca aggtgtttta ggtgaacaag gtagaagaat ccatgaatta acttcattga 720 ttgttaaaag attcaaatta tctccagaag gtattgccat ttatgctgaa agagttgaag 780 aaagaggttt atctgctgct gttcaagctg aagctttgaa agccaaatta ttgtctggtt 840 taccaattag aagagctgct tatggtgttt taagatttgc tatgggtgcc ggtgctaaag 900 gtgttgaagt tgttatctct ggtaaattaa gagctgctag agctaaatct caaaaatatg 960 ctgatggttt tatgattcat tctggtcaac caactagaga tttcattgat attgccatta 1020 gacatgtttt aatgagacaa ggtgttttgg gtatcaaagt taaaattatg aaagatccag 1080 ctgctaatag atttggtcca agagctttac cagatgctgt taaaattgct gaagctaaag 1140 atgaagatga agttattcca gctccaactg ttaaatctta taaacaaact gctgaagatg 1200 aaactgaaac tgatgctcca gttgaagctg aagctgaagt tgaagctact gcttaa

<210> 432

<211> 251

<212> PRT

<213> Candida albicans

<400> 432

Met Val Asn Ala Ile Leu Ser Lys Lys Lys Leu Val Ala Asp Gly

1 5 10 15

Val Phe Tyr Ala Glu Leu Asn Glu Phe Phe Thr Arg Glu Leu Ala Glu 20 25 30

Gln Gly Tyr Ala Gly Val Glu Val Arg Lys Thr Pro Ser Lys Leu Glu 35 40 45

Val Ile Val Lys Ala Ser Asn Thr Gln Gly Val Leu Gly Glu Gln Gly 50 55 60

Arg Arg Ile His Glu Leu Thr Ser Leu Ile Val Lys Arg Phe Lys Leu 65 70 75 80

Ser Pro Glu Gly Ile Ala Ile Tyr Ala Glu Arg Val Glu Glu Arg Gly 85 90 95

Leu Ser Ala Ala Val Gln Ala Glu Ala Leu Lys Ala Lys Leu Leu Ser 100 105 110

Gly Leu Pro Ile Arg Arg Ala Ala Tyr Gly Val Leu Arg Phe Ala Met 115 120 125

Gly Ala Gly Ala Lys Gly Val Glu Val Val Ile Ser Gly Lys Leu Arg 130 135 140

Ala Ala Arg Ala Lys Ser Gln Lys Tyr Ala Asp Gly Phe Met Ile His 145 150 155 160

Ser Gly Gln Pro Thr Arg Asp Phe Ile Asp Ile Ala Ile Arg His Val 165 170 175

Leu Met Arg Gln Gly Val Leu Gly Ile Lys Val Lys Ile Met Lys Asp 180 185 190

Pro Ala Ala Asn Arg Phe Gly Pro Arg Ala Leu Pro Asp Ala Val Lys 195 200 205

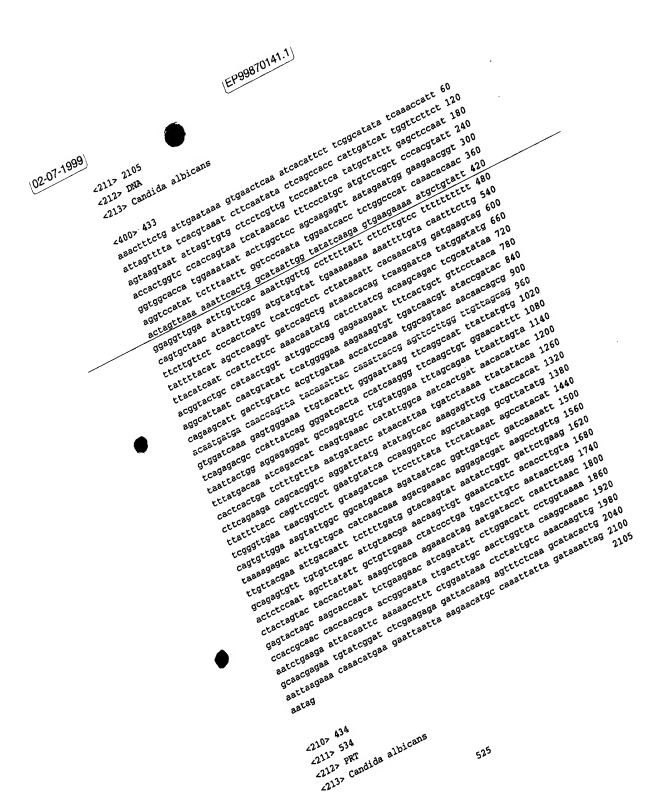
Ile Ala Glu Ala Lys Asp Glu Asp Glu Val Ile Pro Ala Pro Thr Val 210 215 220

Lys Ser Tyr Lys Gln Thr Ala Glu Asp Glu Thr Glu Thr Asp Ala Pro 225 230 235 240

Val Glu Ala Glu Val Glu Ala Thr Ala 245 250

<210> 433

SEOL





<400> 434 Met Tyr Val Leu Lys Lys Lys Asn Phe Val Gln Phe Leu Val Leu Val 10 Leu Pro Leu Ile Ser Ser Leu Ser Tyr Lys Phe Thr Asn Met Asp Glu 20 25 Val Val Phe Tyr Ile Ala Gln Gly Asp Pro Ala Asp Lys His Ser Gln 40 Glu Ser Tyr Gly Tyr Val Thr Ser Ile His Ser Ser Lys Gln Tyr Ala \_\_55\_ \_\_\_\_\_60 Ser Tyr Arg Gln Ala Asp Ser His Ile Asn Gly Thr Ala Ile Thr Gly 70 Ile Gly Pro Gly Glu Arg Ile Phe Thr Ala Val Pro Asn Lys Ala Leu 90 Ile Asn Val Tyr Ser Trp Gly Lys Glu Ser Val Asp Gln Arg Ile Pro 105 Ile Pro Glu Ala Leu Thr Cys Ile Thr Leu Ile Asn His Pro Asn Gly 115 120 Ser Asn Asn Asn Ser Asp Asn Asp Asn Gln Leu Tyr Lys Leu Pro . 130 135 Asn Tyr Arg Val Pro Trp Leu Leu Ala Gly Gly Ser Lys Ser Gly Lys 150 Leu Tyr Ile Trp Glu Leu Ser Ser Gly Asn Leu Leu Cys Val Arg Asp 170 165 Ala His Tyr Gln Gly Ile Thr Thr Ile Lys Gly Ser Ser Cys Gly Thr 180 185

Phe Leu Ile Thr Gly Gly Glu Asp Ala Arg Cys Leu Val Trp Asn Leu 195 200 205

Ala Glu Leu Ile Ser Ile Tyr Asp Lys Ser Asp His Gln Val Lys Pro 210 215 220

Tyr Trp Gln Ile Thr Asp Asn Thr Leu Pro Leu Thr Asp Leu Cys Leu 225 230 235 240

Asn Asp Thr His Asn Ile Asn Asp Leu Lys Leu Tyr Thr Thr Ser Glu 245 250 255

Asp Ser Thr Val Arg Ile Tyr Asp Ile Val Thr Lys Ser Leu Leu Thr 260

Thr Phe Ile Leu Pro Ser Ser Ala Glu Cys Ile Thr Lys Asp Pro Ala 285

Asn Arg Ala Leu Tyr Val Gly Leu Asn Asn Gly Leu Val Arg Ser Ile 290

Pro Leu Tyr Ser Ile Asn Ser His Thr Ser Val Leu Glu Ser Ile Gly 305 310 315 320

Gly Met Asn Lys Ile Ile Thr Val Asp Ala Asp Gln Asn Leu Lys Glu 325 330 335

Thr Phe Val Ala His Gln Gln Lys Thr Lys Thr Gly Asp Asp Lys Pro 340 345 350

Val Val Val Thr Lys Leu Thr Ile Ser Phe Asp Gly Thr Ser Ile Ile 355 360 365

Ser Gly Asp Ser Glu Gly Arg Val Phe Val Ser Asp Ile Val Thr Lys 370 375 380

Gln Val Val Lys Ser Phe Thr Pro Cys Asn Ser Pro Ile Ala Tyr Ile 385 390 395 400

Ala Val Glu Thr Ile Pro Asp Asp Phe Val Asn Asn Leu Ala Thr Ser 405 410 415

Thr Thr Asn Lys Ala Asp Lys Lys His Arg Met Ile Pro Gln Phe 420 425 430

Lys Arg Val Leu Ala Ser Thr Asn Ser Glu Glu His Gln Ile Phe Leu 435 440 445

Asp Ile Pro Gly Lys Thr Thr Ala Thr Thr Asn Ala Thr Gly Asn Ile 450 455 460

Asp Phe Ala Thr Trp Leu Gln Gly Lys Gln Ser Glu Glu Leu Gln Phe 465 470 475 480

Lys Asn Leu Ser Gly Ile Asn Ser Ile Val Lys Gln Val Gly Asn Glu 485 490 495

Asn Val Ser Asp Leu Glu Glu Arg Leu Gln Arg Val Ser Gln Ala Tyr 500 505 510

SEQL

```
Thr Glu Leu Arg Asn Lys His Glu Glu Leu Ile Lys Glu His Ala Lys 515 520 525
```

Leu Leu Asp Lys Leu Glu 530

```
<210> 435
```

<211> 896

<212> DNA

<213> Candida albicans

```
<400> 435
```

tttctgcaag tattgcgtgt gctgcacaaa ctttttatat ccgaaaattt ttggccggga 60 acgaaacggc aagaaaaaac agaaacaata ccaccagcac ggacaaaaag attataagct 120 ttgtgtaata aggttatgtc atcgggtatt acagattgca gggccatctt gtcttcatca 180 gttatagcat ttcaataaaa ataagccaca tatgtgtaca gcgctgagtc tactcaacat 240 gtgtaaatag aataaatcaa ttgacacagt cttttgagat ctgttattct ggcctatagc 300 gttttaggaa attgcggtat tttcttgtct gtttttcttt tatctatttt cgcacgactt 360 ggggtggttt gtgtgacttt tttagcaaat taattttgtc ggtcttcgca gtaaaaataa 420 aaaattcaaa aaaaaaaaca aattgacttt tttttttact ttctttcttt tctatcaaca 480 atactaatca caagccaacc atgaaattca ctactgttgc cactgttttt gctatttcct 540 cattagctgc cgctaaaggt ggtgaaaaag atcacggtaa agcttctact gtcaccaaat 600 atgtcactga aactacccac agatacggtc gttttgacaa aaccagtaga tctaaaaagc 660 caaaggaaac tggtactcac agatacggta aattcaacaa gactccacgt ccagttacca 720 caactgtctt ggtcaaagaa agcgaccttc caaagaaaag agatgctgtt gttgctagag 780 attctaaaaa cgcttcttcc aactctacca cctctagtgg taacaatggt gtcgccactg 840 gtgtcagctt gggtcttgct ggtgtcttag ctgttggtgc tgctttggtc atctaa 896

```
<210> 436
```

<211> 131

<212> PRT

<213> Candida albicans

<400> 436

Met Lys Phe Thr Thr Val Ala Thr Val Phe Ala Ile Ser Ser Leu Ala 1 5 10 15

Ala Ala Lys Gly Gly Glu Lys Asp His Gly Lys Ala Ser Thr Val Thr
20 25 30

Lys Tyr Val Thr Glu Thr Thr His Arg Tyr Gly Arg Phe Asp Lys Thr 35 40 45

Ser Arg Ser Lys Lys Pro Lys Glu Thr Gly Thr His Arg Tyr Gly Lys

50 55 60

Phe Asn Lys Thr Pro Arg Pro Val Thr Thr Thr Val Leu Val Lys Glu 65 70 75 80

Ser Asp Leu Pro Lys Lys Arg Asp Ala Val Val Ala Arg Asp Ser Lys 85 90 95

Asn Ala Ser Ser Asn Ser Thr Thr Ser Ser Gly Asn Asn Gly Val Ala 100 105 110

Thr Gly Val Ser Leu Gly Leu Ala Gly Val Leu Ala Val Gly Ala Ala

115 120 125

Leu Val Ile 130

02-07-1999

<210> 437

<211> 1076

<212> DNA

<213> Candida albicans

<400> 437

tttttgatca ggccttttgt ttttttttt gggtggtgct gtggttcgtt gggtagttgg 60 ctcttgtttc ggttctactt gttccttttt tttattctcc tcgttaatcc taattttgtg 120 taacaaatta attataggga gttggaaaat taaaagcttg aaaagaaaga aagaaagaaa 180 taccaacgtg gaatttctat tacgtaagtc actataactt gcatagaaat ttcaggtttt 240 caatttaaga aagtattaat caactgaatt aagcaattga aacgaattga accagctcag 300 cattlatttt tcgttttctt ttttttcaa gggggtgggt gaaagaaaaa tctaaaaata 360 tataaatact ccacttatct cctctcttcc tctctttctc tctctaactc aatttcaatt 420 atattaaaac aataaaagcc atgtcagcta acgattttta ttcatctggt gatcaatcca 540 attatgatcc aaaaagatcc tcgaatcaag gatcatcatc atcaaatgat gaacaacaag 600 acagagggtt attatctact gtcgccggtg gtgttgctgg tggttatggt ggtcacaaat 660 taggtgaaaa ggcacaacat ggtactttgg gtactgtatt aggtgccatt gggggtgcca 720 ttggtgccaa taaactagaa gatgcttatg aagaccgtaa agaacataaa aaacacgagc 780 aacaatatgg tggtagtggt aaacacgaag gcggaagaca tgaaggtggt tttggtggtg 840 qtaqaccaga tgatcgttat gaaggcgata gaagaaatga taattacggt ggtggttaca 900 atgatagaag agatgacggt tatggtggtg gttacggtgg tggcagacca gacgatagaa 960 gacacgaagg tggtttcggc ggtggcagac cagatgaccg ttttggtggc ggtagaccag 1020 atgaccgttt tggaggtgac agaagagatg atagaagaga tgaccgtaga tggtaa

<210> 438

<211> 191

<212> PRT

SEQL

<213> Candida albicans

<400> 438

Met Ser Ala Asn Asp Phe Tyr Ser Ser Gly Asp Gln Ser Asn Tyr Asp 1 5 10 15

Pro Lys Arg Ser Ser Asn Gln Gly Ser Ser Ser Ser Asn Asp Glu Gln
20 25 30

Gln Asp Arg Gly Leu Leu Ser Thr Val Ala Gly Gly Val Ala Gly Gly
35 40 45

Tyr Gly Gly His Lys Leu Gly Glu Lys Ala Gln His Gly Thr Leu Gly 50 55 60

Thr Val Leu Gly Ala Ile Gly Gly Ala Ile Gly Ala Asn Lys Leu Glu 65 70 75 80

Asp Ala Tyr Glu Asp Arg Lys Glu His Lys Lys His Glu Gln Gln Tyr 85 90 95

Gly Gly Ser Gly Lys His Glu Gly Gly Arg His Glu Gly Gly Phe Gly
100 105 110

Gly Gly Arg Pro Asp Asp Arg Tyr Glu Gly Asp Arg Arg Asn Asp Asn 115 120 125

Tyr Gly Gly Gly Tyr Asn Asp Arg Arg Asp Asp Gly Tyr Gly Gly Gly 130 135 140

Tyr Gly Gly Gly Arg Pro Asp Asp Arg Arg His Glu Gly Gly Phe Gly 145 150 155 160

Gly Gly Arg Pro Asp Asp Arg Phe Gly Gly Gly Arg Pro Asp Asp Arg 165 170 175

Phe Gly Gly Asp Arg Arg Asp Asp Arg Arg Asp Asp Arg Trp
180 185 190

<210> 439

<211> 1745

<212> DNA

<213> Candida albicans

<400> 439

ttggtattga agacaccgaa gacttgttga aagatattga acaagcttta caaaaggctg 60

cttctgtttg aggggatgtt cattagcaat gtatataatt attgtatatt atgacaaaga 120

```
aagaaaaaag aaaaccagaa aagtggttta tacaggaata ttttaataga aatatcgctt 180
atattgtgat aaaaaatttg aaagacaatc cgaatgtagt gcttgtctta ttctgcttgg 240
gaatactgta gtattagcat caattgagga aattccagat agctaacggt tttgcgatta 300
cgaatttcgc aaccaaataa atatgtgaca aggaatacac tactgatcaa ggttattctt 360
agtacaatgg aaaaaaaaa aaagaagcaa acaaaaaaac gagaaattaa tgaacacgac 420
ttcacttcta caacctactg ggaaaaaaaaa ggcagagagt tattgaaaaa ggatcatatc 480
aagttettat tgtatattat atgaagtttt etgttttagt attacttgee agttacttag 540
ttggtgtgaa ttctctgatt gttgatactt cagaggaatt aatttgtcca gatccagaaa 600
accetttaga ttgttateca aaattgtttg ttecaacaaa egagtggcaa accattaaac 660
caggtcaaga tataccacct gggttacacg ttagattaaa tatagatacg ttggaaaaag 720
aggccaaget aatgagtget gaegaaaaag aegageeagt teaagaagta gttgttggtg 780
gcgaattgca ggatcattcg agggaagcca tcactgagaa tctacaaaag ttgcatgagc 840
tgaaacatcc tgaagtaaaa caggagcacg ctcatcgtac aaaggttagc cagggagatt 900
tgagtaattt tgacgcagct tgtctggaaa ttgagagttt caagccacat gagagtgatg 960
tggaaaggtt gcatttggca ctagatactt tagaggaatt aagtcatgat atcgaatttg 1020
gggtgaaatt gacctcagac aaagccatat ttcagagttt tgtcaacatt gccaatggtg 1080
cttctgatcc aaaaataacc gaaaaggtat atcgtgtaat ggggtctagt ttgagaaata 1140
atcctgaagc gattagtaat atcttgacca acttcgacaa gagctatgtg gataatttgt 1200
ttgagcaatt agcgaatgaa aatgatgttc tacaaaagag gattttgggt ataattcaag 1260
etttagteca aaatageeat tttgeaagae aatattttte atttgaceae agtteegggt 1320
taaatgattt aatagcgatt tttcccaaac ttggtccaaa ctcaaagtcc agggcaagta 1380
acattttaga ggatttacaa ttgttcccag taacaaacga tagaagatca cttgaagatc 1440
aagateetga ateacaggtt teaaaattta tteagaatte etttgttgga aataaaettg 1500
acgagaagaa tttcaagtct tattttgatc aactagtaaa tttgcatcag ctgaataaga 1560
gtttgcgacc aagtggtgac tttctcaatt ggttagctga agaagtggag tcgcgtaaag 1620
agaataaaaa aagagacgat tattcacaag aagacaaaga ctttgatgag tacatgttgc 1680
gagcacgtca tgaagtattt ggcaatccaa tgggattaag aaaggcaatt gccgacgagt 1740
tgtag
                                                                  1745
```

```
<210> 440
```

<211> 414

<212> PRT

<213> Candida albicans

<400> 440

Met Lys Phe Ser Val Leu Val Leu Leu Ala Ser Tyr Leu Val Gly Val
1 5 10 15

Asn Ser Ser Ile Val Asp Thr Ser Glu Glu Leu Ile Cys Pro Asp Pro 20 25 30

Glu Asn Pro Leu Asp Cys Tyr Pro Lys Leu Phe Val Pro Thr Asn Glu 35 40 45

Trp Gln Thr Ile Lys Pro Gly Gln Asp Ile Pro Pro Gly Leu His Val
50 55 60

65	Dea	Non	110	nsp	70	Deu	Giu	D, 3	014	75	2,5	200		501	80
Asp	Glu	Lys	Asp	Glu 85	Pro	Val	Gln	Glu	Val 90	Val	Val	Gly	Gly	Glu 95	Leu
Gln	Asp	His	Ser 100	Arg	Glu	Ala	Ile	Thr 105	Glu	Asn	Leu	Gln	Lys 110	Leu	His
Glu	Ser	Lys 115		Pro	Glu		Lys 120		Glu	His		His 125	Arg	Thr	Lys
Val	Ser 130	Gln	Gly	Asp	Leu	Ser 135	Asn	Phe	Asp	Ala	Ala 140	Суѕ	Ser	Glu	Ile
Glu	Ser	Phe	Lys	Pro	His	Glu	Ser	Asp	Val	Glu	Arg	Leu	His	Leu	Ala
145					150					155					160
Leu	Asp	Thr	Leu	Glu 165	Glu	Leu	Ser	His	Asp 170	Ile	Glu	Phe	Gly	Val 175	Lys
Leu	Thr	Ser	Asp 180	Lys	Ala	Ile	Phe	Gln 185	Ser	Phe	Val	Asn	Ile 190	Ala	Asn
Gly	Ala	Ser 195	Asp	Pro	Lys	Ile	Thr 200	Glu	Lys	Val	Tyr	Arg 205	Val	Met	Gly
Ser	Ser 210	Leu	Arg	Asn	Asn	Pro 215	Glu	Ala	Ile	Ser	Asn 220	Ile	Leu	Thr	Asn
Phe 225	Asp	Lys	Ser	Tyr	Val 230	Asp	Asn	Leu	Phe	Glu 235	Gln	Leu	Ala	Asn	Glu 240
Asn	Asp	Val	Leu	Gln 245	Lys	Arg	Ile	Leu	Gly 250	Ile	Ile	Gln	Ala	Leu 255	Val
Gln	Asn	Ser	His 260	Phe	Ala	Arg	Gln	Tyr 265	Phe	Ser	Phe	Asp	His 270	Ser	Ser
Gly	Leu	Asn 275	Asp	Leu	Ile	Ala	Ile 280	Phe	Pro	Lys	Leu	Gly 285	Pro	Asn	Ser
Lys	Ser 290	Arg	Ala	Ser	Asn	Ile 295	Leu	Glu	Asp	Leu	Gln 300	Leu	Phe	Pro	Val
Thr 305	Asn	Asp	Arg	Arg	Ser 310	Leu	Glu	qaA	Gln	Asp 315	Pro	Glu	Ser	Gln	Val 320

 Ser Lys
 Phe Ile Gln Asn Ser Phe Val Gly Asn Lys
 Leu Asp Glu Lys

 325
 325

 Asn Phe Lys
 Ser Tyr Phe Asp Gln Leu Val Asn Leu His Gln Ser Asn

 340
 345

 Lys
 Ser Leu Arg Pro Ser Gly Asp Phe Leu Asn Trp Leu Ala Glu Glu

 355
 360

Asp Lys Asp Phe Asp Glu Tyr Met Leu Arg Ala Arg His Glu Val Phe 385 390 395 400

Val Glu Ser Arg Lys Glu Asn Lys Lys Arg Asp Asp Tyr Ser Gln Glu

Gly Asn Pro Met Gly Leu Arg Lys Ala Ile Ala Asp Glu Leu 405 410

375

<210> 441 <211> 1244 <212> DNA <213> Candida albicans

#### <400> 441

atgatttttt ccttgatttt agagctgtta attatacctt ttttgttccc aaataagcgc 60 taaaaatgat tgaaattaaa aatactaact attccaggtc aaagtaaaca ctataaaaca 120 atatatttca ataacaggag cagtaattag cttcaacatc aaggtactct tttatttttc 180 taccaaaaac acatctgaag tagctcttat ccatagatcc aaatatttta accttttttt 240 tttcctactc tcatctactt ttttttgcaa cacttactgc tcacaacgcc aatgaccata 300 ccattaattt caataatcaa atcaagagct tatttgtatc ctctcaaggt atgttaatgt 360 attaacaaca ccgattctat ttcaccaact aacacgacag aaaggggttg tactattttg 420 taacacatcc caccgtttgg cccttttaca taaccatatt gatacctcaa ttggtcctta 480 cactagteat ttacctgatt atgttttcat tgttcttccc acctcaaget attgtatata 540 cattgttaat gggaccatta ggggtgattg gtgcgtggta tagtttgatc ctgcaagcga 600 gcacattgtc tatatttgtt gtcacaatat ccttaatgcc tcacatacaa cgagtggcat 660 atgatgcaat attgagtaga gagtgtgcaa atgatgtggt actaatggga aaacttagga 720 ggtatagaaa actacccatt agagtaaggg ccagagaata ccttaaggcc attccagatt 780 tttcgatctt tcccttctct ttgctcaagc tacttgtgtt tttcggtatt tacttcatac 840 cctttgtagg tccaatcatt gtattgtttt tccaatcttc caagcgtgga ctaaaggcac 900 atgcaagata ctttaagttg aaagggtttc tgcgtagtga cataagaaca atccacaagc 960 taaacagacc agcatatatg gggtacggag tggttgcgct ttggctcgag ctgtttccat 1020 ttatcaatat gttttttatg ttcaccaata ctttgggagc tgctttgtgg gcagttgata 1080 ttgaacaaca agagaaggcc gtcacagaga atgtggccgc agctactacc accgccacag 1140 atacgaatag cgtcaatcaa caaggtctag ttatacccgt acacaatgaa ccagcaacta 1200 atatacctga ggctacccca aaaactgcta caaataccat ctaa 1244 <210> 442

<211> 247

<212> PRT

<213> Candida albicans

<400> 442

Met Phe Ser Leu Phe Phe Pro Pro Gln Ala Ile Val Tyr Thr Leu Leu 1 5 10 15

Met Gly Pro Leu Gly Val Ile Gly Ala Trp Tyr Ser Leu Ile Ser Gln
20 25 30

Ala Ser Thr Leu Ser Ile Phe Val Val Thr Ile Ser Leu Met Pro His 35 40 45

Ile Gln Arg Val Ala Tyr Asp Ala Ile Leu Ser Arg Glu Cys Ala Asn 50 55 60

Asp Val Val Leu Met Gly Lys Leu Arg Arg Tyr Arg Lys Leu Pro Ile 65 70 75 80

Arg Val Arg Ala Arg Glu Tyr Leu Lys Ala Ile Pro Asp Phe Ser Ile 85 90 95

Phe Pro Phe Ser Leu Leu Lys Leu Leu Val Phe Phe Gly Ile Tyr Phe 100 105 110

Ile Pro Phe Val Gly Pro Ile Ile Val Leu Phe Phe Gln Ser Ser Lys
115 120 125

Arg Gly Leu Lys Ala His Ala Arg Tyr Phe Lys Leu Lys Gly Phe Ser 130 135 140

Arg Ser Asp Ile Arg Thr Ile His Lys Leu Asn Arg Pro Ala Tyr Met 145 150 155 160

Gly Tyr Gly Val Val Ala Leu Trp Leu Glu Ser Phe Pro Phe Ile Asn 165 170 175

Met Phe Phe Met Phe Thr Asn Thr Leu Gly Ala Ala Leu Trp Ala Val 180 185 190

Asp Ile Glu Gln Gln Glu Lys Ala Val Thr Glu Asn Val Ala Ala Ala 195 200 205

Thr Thr Thr Ala Thr Asp Thr Asn Ser Val Asn Gln Gln Gly Leu Val

210 215 220

Ile Pro Val His Asn Glu Pro Ala Thr Asn Ile Pro Glu Ala Thr Pro 225 230 235 240

Lys Thr Ala Thr Asn Thr Ile 245

<210> 443

<211> 2270

<212> DNA

<213> Candida albicans

<400> 443

gcattttact tatttagata ttacttcata ttgcttttat ttagattgat tttgtttaac 60 tgtattatac aatatttaaa ttatagtaat catcctataa atttcaaagt caaaagacag 180 atcttaaggt ctaattaata actetetatg geettetgtg teaaattgtt gtegtttgat 240 ctgatgtata gaactttcta gcaaaaaaaa aaagaaagaa tttttttttc ttccattttt 360 caaatttgag agatcgaaat aattttcttg aatttattaa aagggaaccc cttcccgaaa 420 aatccaaaac caaaacttcc acccaaatat caaataacta acttatcatt ccaacagata 480 atattcccac ttcaataaca atgacaacag ctgacgaata caaagcagaa ggtaacaaat 540 attttgctgc taaagatttt gaaaaggcga ttgaagcatt cactaaagca attgaagcat 600 cacctgaacc aaaccatgtt ctttattcaa atcgttctgg atcttatgcc tctttaaaag 660 attttaacaa cgcattaaaa gatgctcaag aatgtgtcaa gatcaatcct agttgggcca 720 aagggtataa tagaattgct ggggctgaat ttggtttagg taattttgat caagccaaat 780 ccaattatga aaaatgtttg gagttggatc caaataatgc catggctaaa gaaggtttaa 840 aatcagttga atctgcttta tcatctggtg gtggtgatga caaggattta ggatttggta 900 aaattttaaa tgatcctaat ctttatacta aattgaaaaa taatcctaaa acaagtgaat 960 ttatgaatga tcctcaattt gttgctaaac ttgaacgtct taaaactaat ccacaattgg 1020 gtaatcctga tatgtttagt gatccaagat tattgacggc ttttgctgct ttaatgggta 1080 ttgacatgga tttaccaaat atgggattca ctgctccaaa cgaatcacaa tccaatgcat 1140 cagaaccaaa actggaacca aaatcagtac cagaatctaa accagaacca aaagcagaac 1200 aaaaggaaga agaatcaacc tcagccaaag atgaagacac tccaatgact gatgcccaag 1260 acgacactaa tgataatgat gccaaaaccc aagctgacaa tgctaaagct gaaggtaatg 1320 ccttatacaa gaaacgtcaa tttgatgaag caattgccgc ctataataag gcttgggaat 1380 tacataagga tatcacttat ttaaacaatc gtgctgctgc cgaatatgaa aaaggtgatt 1440 atgatgctgc tattgctaca tgtgaaaagg ccattgatga aggtagagac atgagagctg 1500 attataaatt gattgctaaa tcatttgcta gattaggtaa tatttatttg aaaaaagatg 1560 aattacccga agcagtgaaa aattttgaaa aatctttaac tgaacatcgt acccctgatg 1620 ttttaaataa attaagatca actcaacgtg aaattaaaac tagagaatta aatgcttata 1680 tagatccaga aaaggctgaa gaagcaagat tacaaggtaa agaatatttc accaaaggag 1740 attggccaaa tgccgttaag gcttatactg aaatgattaa aagagcacca gaagatgcta 1800 gaggatattc taatcgtgct gctgcattgg caaaattgtt atcatttcct gatgctatac 1860 aagattgtaa taaagccatt gaaaaagatc caaatttcat tagagcttat attagaaaag 1920

<210> 444

<211> 589

<212> PRT

<213> Candida albicans

<400> 444

Met Thr Thr Ala Asp Glu Tyr Lys Ala Glu Gly Asn Lys Tyr Phe Ala 1 5 10 15

Ala Lys Asp Phe Glu Lys Ala Ile Glu Ala Phe Thr Lys Ala Ile Glu 20 25 30

Ala Ser Pro Glu Pro Asn His Val Leu Tyr Ser Asn Arg Ser Gly Ser
35 40 45

Tyr Ala Ser Leu Lys Asp Phe Asn Asn Ala Leu Lys Asp Ala Gln Glu 50 55 60

Cys Val Lys Ile Asn Pro Ser Trp Ala Lys Gly Tyr Asn Arg Ile Ala 65 70 75 80

Gly Ala Glu Phe Gly Leu Gly Asn Phe Asp Gln Ala Lys Ser Asn Tyr 85 90 95

Glu Lys Cys Leu Glu Leu Asp Pro Asn Asn Ala Met Ala Lys Glu Gly
100 105 110

Leu Lys Ser Val Glu Ser Ala Leu Ser Ser Gly Gly Asp Asp Lys
115 120 125

Asp Leu Gly Phe Gly Lys Ile Leu Asn Asp Pro Asn Leu Tyr Thr Lys 130 135 140

Leu Lys Asn Asn Pro Lys Thr Ser Glu Phe Met Asn Asp Pro Gln Phe 145 150 155 160

Val Ala Lys Leu Glu Arg Leu Lys Thr Asn Pro Gln Leu Gly Asn Pro
165 170 175

Asp Met Phe Ser Asp Pro Arg Leu Leu Thr Ala Phe Ala Ala Leu Met

185

190

Gly Ile Asp Met Asp Leu Pro Asn Met Gly Phe Thr Ala Pro Asn Glu 195 200 205

Ser Gln Ser Asn Ala Ser Glu Pro Lys Ser Glu Pro Lys Ser Val Pro 210 215 220

Glu Ser Lys Pro Glu Pro Lys Ala Glu Glu Lys Glu Glu Glu Ser Thr 225 230 235 240

Ser Ala Lys Asp Glu Asp Thr Pro Met Thr Asp Ala Gln Asp Asp Thr

245
250
255

Asn Asp Asn Asp Ala Lys Thr Gln Ala Asp Asn Ala Lys Ala Glu Gly 260 265 270

Asn Ala Leu Tyr Lys Lys Arg Gln Phe Asp Glu Ala Ile Ala Ala Tyr 275 280 285

Asn Lys Ala Trp Glu Leu His Lys Asp Ile Thr Tyr Leu Asn Asn Arg 290 295 300

Ala Ala Glu Tyr Glu Lys Gly Asp Tyr Asp Ala Ala Ile Ala Thr 305 310 315 320

Cys Glu Lys Ala Ile Asp Glu Gly Arg Asp Met Arg Ala Asp Tyr Lys 325 330 335

Leu Ile Ala Lys Ser Phe Ala Arg Leu Gly Asn Ile Tyr Leu Lys Lys 340 345 350

Asp Glu Leu Pro Glu Ala Val Lys Asn Phe Glu Lys Ser Leu Thr Glu 355 360 365

His Arg Thr Pro Asp Val Leu Asn Lys Leu Arg Ser Thr Gln Arg Glu 370 375 380

Ile Lys Thr Arg Glu Leu Asn Ala Tyr Ile Asp Pro Glu Lys Ala Glu
385 390 395 400

Glu Ala Arg Leu Gln Gly Lys Glu Tyr Phe Thr Lys Gly Asp Trp Pro
405 410 415

Asn Ala Val Lys Ala Tyr Thr Glu Met Ile Lys Arg Ala Pro Glu Asp 420 425 430

Ala Arg Gly Tyr Ser Asn Arg Ala Ala Ala Leu Ala Lys Leu Leu Ser

440

445

Phe Pro Asp Ala Ile Gln Asp Cys Asn Lys Ala Ile Glu Lys Asp Pro 450 455 460

Asn Phe Ile Arg Ala Tyr Ile Arg Lys Ala Asn Ala Gln Leu Ala Met 465 470 475 480

Lys Glu Tyr Ser His Val Met Asp Thr Leu Thr Glu Ala Arg Thr Lys
485 490 495

Asp Val Glu Leu Gly Gly Lys Ser Ile His Glu Ile Asp Glu Leu Met
500 505 510

Asn Lys Ala Thr Tyr Gln Arg Phe Gln Ala Ile Glu Gly Glu Thr Pro 515 520 525

Glu Gln Thr Met Glu Arg Val Ser Lys Asp Pro Glu Ile Val Gln Ile 530 535 540

Leu Gln Asp Pro Val Met Gln Gly Ile Leu Ala Gln Ala Arg Glu Asn 545 550 555 560

Pro Ala Ala Leu Gln Asp His Met Lys Asn Pro Glu Val Tyr Lys Lys 565 570 575

Ile Asn Met Leu Ile Ala Ala Gly Val Ile Arg Thr Arg
580 585

<210> 445

<211> 1019

<212> DNA

<213> Candida albicans

### <400> 445

ccactaacaa actttttt gactatacac cactgaaaaa aaaaaaaaa ttttgtaaa 60 aaatcttaag aattcagtt tagaaggtat tcaacaacaa ccgaagagta tgttcaacat 120 cgttataata gaatagaata agagcatgac aacaaaggga tacaagcttg aaaaaaagaaa 180 aagggtggga tatcttaaaa ttattaaaga gttttttta taacatgtca ttgagattga 240 gattgggaat actgaattcg attttaaagt cattggatgg gagagttaat tattcgttt 300 attattagga ttaccaatga atagtaatga agtgatggag agatagaatg aaagtattca 360 gaagagcatc aagtccctta taagtttgtg agacataata tgtctacccc cttgtcaact 420 tgtcataaat ttattgctc gtcctttaaa agaaatgaat aaaaagattt actaacttaa 480 tttcaattat ttatagaaag atgtctagat taaacgaata tcaagttatt ggtcgtaatt 540 ccgttgttgc caaatcaaga tattggtatt tcttgcaaaa attgcataaa gttaaaaaag 660

cttctggtga aattgtatct gtcaacatta tttctgaagc taaaccaact aaagttaaaa 720 cttttggtat ttggttaaga tatgaatcca gatctggtat tcataacatg tacaaagaat 780 acagagatgt tactagagtt ggtgctgttg aaaccatgta ccaagattta gctgctagac 840 acagagctag atttagaagt atccatattt tgaaagttgt tgaattagaa aaaactgatg 900 atgttaaaag acaatacgtt aaacaatttt tgactaaaga tttgaaattc ccattaccac 960 acagagtcca aaaatctaag aaattgttcc aagctactgc tccaaccact ttctactaa 1019

<210> 446

<211> 172

<212> PRT

<213> Candida albicans

<400> 446

Met Ser Arg Leu Asn Glu Tyr Gln Val Ile Gly Arg Asn Leu Pro Thr
1 5 10 15

Glu Ser Val Pro Glu Pro Lys Leu Phe Arg Met Arg Ile Phe Ala Pro 20 25 30

Asn Thr Val Val Ala Lys Ser Arg Tyr Trp Tyr Phe Leu Gln Lys Leu 35 40 45

His Lys Val Lys Lys Ala Ser Gly Glu Ile Val Ser Val Asn Ile Ile 50 55 60

Ser Glu Ala Lys Pro Thr Lys Val Lys Thr Phe Gly Ile Trp Leu Arg
65 70 75 80

Tyr Glu Ser Arg Ser Gly Ile His Asn Met Tyr Lys Glu Tyr Arg Asp
85 90 95

Val Thr Arg Val Gly Ala Val Glu Thr Met Tyr Gln Asp Leu Ala Ala 100 105 110

Arg His Arg Ala Arg Phe Arg Ser Ile His Ile Leu Lys Val Val Glu 115 120 125

Leu Glu Lys Thr Asp Asp Val Lys Arg Gln Tyr Val Lys Gln Phe Leu 130 135 140

Thr Lys Asp Leu Lys Phe Pro Leu Pro His Arg Val Gln Lys Ser Lys 145 150 155 160

Lys Leu Phe Gln Ala Thr Ala Pro Thr Thr Phe Tyr 165 170

<210> 447

```
<211> 932
<212> DNA
<213> Candida albicans
<400> 447
tgtttacttt ttctgtagtt ttaaagtttc ctaatttaac ttccaaaagt ttcattaaca 60
acaatattaa catacctttg tcacaagcaa ttatattgaa gttttttgat acaagtgtgt 120
tgtttttttg tgtacatgtg agatatataa ttgtgtatat acagtcacgt gaatagagca 180
gaaaaattac gaagtagaaa tattggtagc gcgttagggc tatagcccta tttagtttgt 240
gcaccacaeg acttacaatt ttttttttt tetttettag aateettgag gcactgacae 300
tgtactctct ctctctctc ctctctctcg taggtagtga aaaatttcca ctagtcttcc 360
cataacccac ctaggttctt tcttttggaa accactgagc agtaaatcaa tttacttgac 420
gaagaagtct atacataaat ataaacttgt cccctcccc cccttttttt ttaactaact 480
aagaagaaaa aattaaaaaa atgtctgacg ttgaacaaga acaaattgtt gaagaagttg 540
ttgttgaaga acaatccggt gccatcacca ttgaagatgc tttaaaagtt gttttaagaa 600
cttctttagt ccatgatggt ttagctagag gtttaagaga agcttctaaa gctttatcta 660
aaagagaagc tcaattatgt gttttgtgtg actctgttac tgaagaatca atcatcaaat 720
tggttgaagc tttatgtaat gaaccagaag aaaaaatccc attgattaaa gtttccgatg 780
ctaaattatt gggtgaatgg gctggtttat gtcaattaga tagagatggt aatgctagaa 840
aagttgttgg tgcctcttgt gttgttgtca aaaactgggg tgctgattct gatgaaagaa 900
acatcttgtt ggaacacttt tctcaacaat aa
                                                                   932
<210> 448
<211> 143
<212> PRT
<213> Candida albicans
<400> 448
Met Ser Asp Val Glu Gln Glu Gln Ile Val Glu Val Val Val Glu
                                     10
Glu Gln Ser Gly Ala Ile Thr Ile Glu Asp Ala Leu Lys Val Val Leu
             20
                                 25
Arg Thr Ser Leu Val His Asp Gly Leu Ala Arg Gly Leu Arg Glu Ala
                             40
Ser Lys Ala Leu Ser Lys Arg Glu Ala Gln Leu Cys Val Leu Cys Asp
     50
                         55
                                             60
Ser Val Thr Glu Glu Ser Ile Ile Lys Leu Val Glu Ala Leu Cys Asn
 65
                     70
                                         75
                                                             80
```

540

85

Glu Pro Glu Glu Lys Ile Pro Leu Ile Lys Val Ser Asp Ala Lys Leu

90

```
Leu Gly Glu Trp Ala Gly Leu Cys Gln Leu Asp Arg Asp Gly Asn Ala
100 105 110
```

Arg Lys Val Val Gly Ala Ser Cys Val Val Val Lys Asn Trp Gly Ala 115 120 125

Asp Ser Asp Glu Arg Asn Ile Leu Leu Glu His Phe Ser Gln Gln 130 135 140

<210> 449

<211> 881

<212> DNA

<213> Candida albicans

### <400> 449

aaaatttcca atcttgaatt tcatcttcaa cgtcataaac ttgttctgtt tgaaatttat 60 cgtcctcatt atcgttgtca ttttcattgt cattgtcaat attacggtta cgattttgtt 120 ttccctttga agtctttgac tctagctgtt catctgtcat tacccttggc ttggtttgtc 180 ttactgttga gcgatgagac aaactttttg attgattgca actgaacaaa aaaaaatacg 240 acagacacac acacacaca acacacaca ttttcagctc cttccaagtc gtgttttttt 300 ggaagaaaaa aaaaacaact tggccctaaa aactctatgc tctaaccgac aacattagat 360 tactttgata actcacaacc tttaattaac actatctaca aaatatgaca agtacaccaa 420 taacgtacaa gacactagta tgaaaggcaa gcacaaactt gcaaacaaaa aacccctaac 480 ttttattata gttttgtttg atgttatttt gttttatact aactaaaatc tgttttttt 540 ttttttcaaa ggccgattct atatttaatg atctaatcaa caatatcatt aaacaacata 600 cattaaccag tttaaccaat attaaagatc attcctcatt attaaattca tctaatagca 660 atactaatag caataccaac ggtacaattg ccagtaatgg tggaaatggg actacaagtg 720 atgaaaataa tgaaattgaa aattcaacaa ttcaagataa atcaaaatta aaacaattag 780 aaacttcaag gtatttccga tgtcttaatt gtggtagaaa tattgccggt ggaagatttg 840 catctcatat aagtaagtgt ttagaacgga aacggaaatg a 881

<210> 450

<211> 126

<212> PRT

<213> Candida albicans

### <400> 450

Met Leu Phe Cys Phe Ile Leu Thr Lys Ile Cys Phe Phe Phe Ser
1 5 10 15

Lys Ala Asp Ser Ile Phe Asn Asp Leu Ile Asn Asn Ile Ile Lys Gln
20 25 30

His Thr Leu Thr Ser Leu Thr Asn Ile Lys Asp His Ser Ser Leu Leu

40

45

Asn Ser Ser Asn Ser Asn Thr Asn Ser Asn Thr Asn Gly Thr Ile Ala 50 55 60

Ser Asn Gly Gly Asn Gly Thr Thr Ser Asp Glu Asn Asn Glu Ile Glu 65 70 75 80

Asn Ser Thr Ile Gln Asp Lys Ser Lys Leu Lys Gln Leu Glu Thr Ser 85 90 95

Arg Tyr Phe Arg Cys Leu Asn Cys Gly Arg Asn Ile Ala Gly Gly Arg
100 105 110

Phe Ala Ser His Ile Ser Lys Cys Leu Glu Arg Lys Arg Lys 115 120 125

<210> 451

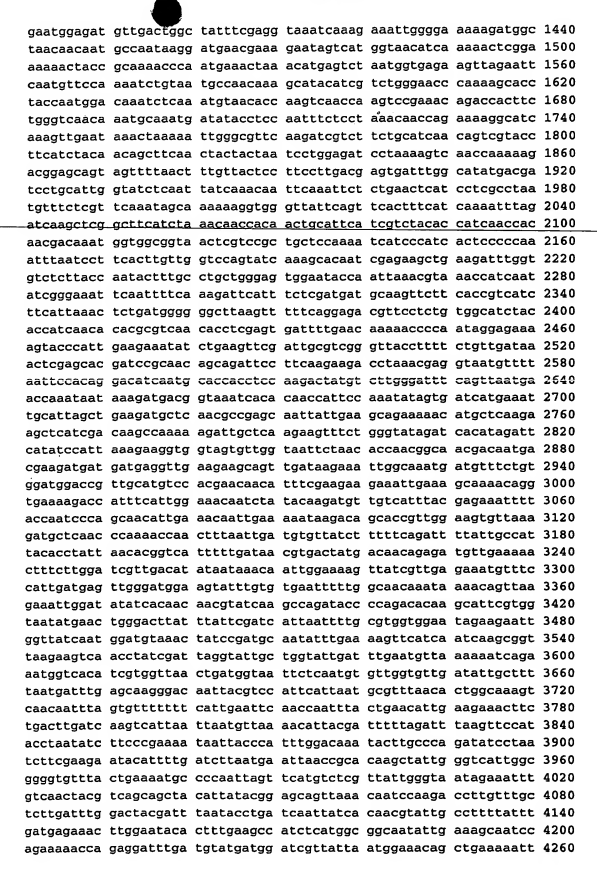
<211> 5344

<212> DNA

<213> Candida albicans

#### <400> 451

ctctagaagt aggacatcgt atagtgtata aacactcaat aagtaatgaa gaaacacgtt 60 ttgttgtgca atgttagctg gcgagctcaa tattgggtct ctccgccgtc tgtctgcctt 120 gcattctact ttttttctcc ttatggaaaa cattagtact agtagtggtg gtagtagttg 180 ctgttgtttg cactttgcac gtagttgttt tcccctcatt ctttcatctg attattctgt 240 ctttgttaac tgcatacaaa gggagggaa gaagaaacaa caaaagggga atttgaatat 300 acgtcaatct tttaatccta ctaccacggg ggggggggg gtcatacttc ttggtgtaaa 360 taatgtatgg agattgaggt tattagactt ttagaaagag gtcttggttg agtaaggcgg 420 gatagcacaa ataatgcgtg tctaattggt cagagaataa tatgctttgg ggaacaatag 480 aaagatgtag cgagagaaat agaattncaa gggtaaggat gattaacttt ttttttttt 540 aatgcttgga gtactttgtt gttggaaaga gtattgcaga agtaatagtt taataaaaga 600 aaagtatata actttagtaa tcggagaaca attgaaatca atattttgaa aatatagttt 660 tatagagtaa cctgggtcga ggtgaacccg aattcaatat tggttttcgg ttgttatatg 720 ctacataacc atcctcatct tgaatgaaac aaggataata gaaatgagta tattaaacaa 780 acaacattat gttgttactg ttgtattctg ttttttggtc attgccatgg taatttcata 840 tttagtaaca atttagtgtt catcgtaagt ctctccttct ctgggagatg ttctctctct 900 ctttctctct cttgttgttg ttgttgttaa ttgttaattg ttgattattg attgttcccc 960 aattaaaatt ccgataaact tgaacaataa attgtcatca taaatttttt tttagaaaat 1020 cactccataa attaaattta tttatatatt aaattttaaa tcggaacttc cgtcaattgg 1080 cctacttctc ttttattttc ttagtttcat aatttttgac agataatact tgaaactggt 1200 taattttaaa tcaacttcgt gattttttc cggaaatcta ataatatcaa tgaccagtaa 1260 tctgccacca cttggttcta caactaacga tcagagacta ccccaaagcg gagtttcatc 1320 catacccaca aataaattgc ctttacctaa tgccaatgaa gattttgcca cgggtgtgtc 1380



```
attagttgaa atagaaaagg gtaagaaaga ggatatcaaa atgcaaagaa ttatatccga 4320
ttcagtattg gaaagaacaa gatcgattcg taaggatatt cacaaaacca ttgatacatt 4380
attcgaacaa agaaatttag gtaaattatc atttgaaggt aaagagaatt tagttcgatt 4440
ttgtttatta gattcgtctt tagaaaaatt ggttgttatg gttgaggaac atgccaacgg 4500
attattatta acaccaacga cctccacgga cgatctcaga agtagagcca tgtcgccatc 4560
ggtcactgtt gatacaatcc atgaaagtgc aaatgagttg attactgctg gaccaatttt 4620
atcaccacat gtcaatagga aagcagaaca aagctcgtat ttcccagtgt ttgccaataa 4680
tgataatttg acccctcatc aagttgtcgt tgagtcaaat gatgaaggta gagatgttcc 4740
aatagataaa atgacaggac gaccagtttt gattcgatca attagtcaaa cttctgtgca 4800
tgcaaaagag caagaaattg aagaagggga gcttcataaa tttggattct ttattcaaca 4860
aaaagagaga caaaaacaac aacaacaaca acaacaacaa cagaactcac accaccagca 4920
ccaaccggcc cagctgatcc aacaagaaaa ccagctgceg ctgccacaac aaggaaaata 4980
tgaagattta ccgatattaa atacattacc gtcaggacca gagttgagag atgctataat 5040
ggcagctaag ggagtagcaa atgttactga attaattgat cgaattaata atcatcgtgt 5100
taaaatcgat gcaccatcga caaaacacca tcatgaattg aacaaaccaa attctgacaa 5160
agtagttgag gatgaagttg aagtttctga taatgcctct attgattcta ctaatggtga 5220
cgatttacat caacttggtg acggtaaaca taatggtaat ggtacggttg atcccatggt 5280
tagtgaagtt tatgacaagt tgttaaatga tgctgaacga gtcagactga atagagatat 5340
                                                                  5344
ataa
```

<210> 452

<211> 1364

<212> PRT

<213> Candida albicans

<400> 452

Met Thr Ser Asn Ser Pro Pro Leu Gly Ser Thr Thr Asn Asp Gln Arg

1 5 10 15

Leu Pro Gln Ser Gly Val Ser Ser Ile Pro Thr Asn Lys Leu Pro Leu
20 25 30

Pro Asn Ala Asn Glu Asp Phe Ala Thr Gly Val Ser Asn Gly Asp Val
35 40 45

Asp Trp Leu Phe Arg Gly Lys Ser Lys Lys Leu Gly Lys Lys Met Ala 50 55 60

Asn Asn Asn Ala Asn Lys Asp Glu Arg Lys Asn Ser His Gly Asn Ile
65 70 75 80

Lys Asn Ser Glu Lys Thr Thr Ala Lys Pro Asn Glu Thr Lys His Glu
85 90 95

Ser Asn Gly Glu Lys Leu Glu Phe Asn Val Pro Lys Ser Val Met Pro 100 105 110 Thr Lys His The Ser Ser Gly Asn Pro Lys Ala Pro Thr Asn Gly Gln Ile Ser Asn Val Thr Pro Ser Gln Pro Ser Pro Lys Gln Thr Thr Ser Gly Ser Thr Asn Ala Asn Asp Ile Pro Pro Ile Ser Pro Lys Gln Pro Glu Lys Ala Ser Lys Leu Asn Lys Leu Lys Ile Gly Arg Ser Arg Ser Ser Ser Ala Ser Thr Val Val Pro Ser Ser Thr Thr Ala Ser Thr Thr Thr Asn Pro Gly Asp Pro Lys Ser Gln Pro Lys Arg Arg Ser Ser Ser Phe Asn Phe Val Thr Pro Ser Leu Thr Ser Asp Leu Ala Tyr Asp Asp Pro Ala Leu Val Ser Gln Leu Ser Asn Asn Ser Asn Ser Ser Asn Ser Ser Ser Pro Asn Val Ser Arg Ser Asn Ser Lys Lys Gly Gly Leu Phe Ser Ser Leu Ser Ser Lys Phe Arg Ser Ser Ser Ala Ser Ser Lys Gln Pro Gln Ser His Ser Ser Ser Thr Pro Ser Thr Thr Thr Thr Asn Gly

Gly Gly Asn Ser Ser Ala Ala Pro Lys Ser Ser His His Ser Pro Lys 

Phe Asn Pro Ser Leu Val Gly Pro Val Ser Lys His Asn Arg Glu Ala 

Glu Asp Leu Val Ser Leu Thr Asn Thr Leu Pro Ala Gly Ser Gly Ile 

Pro Ile Lys Arg Lys Pro Ser Ile Ser Gly Asn Ser Ile Phe Lys Asp 

Ser Phe Leu Asp Asp Ala Ser Ser Ser Pro Ser Ser Ser Leu Asn Ser 

Asp Gly Gly Leu Lys Phe Phe Arg Arg Ser Ser Val Ala Ser Thr Pro Ser Thr His Ala Ser Thr Pro Arg Val Ile Leu Asn Lys Asn Pro Asn Arg Arg Lys Val Pro Ile Glu Glu Ile Ser Glu Val Arg Leu Arg Arg Val Thr Phe Ser Val Asp Lys Leu Glu His Asp Pro Gln Gln Gln Ile Pro Ser Arg Arg Pro Lys Arg Gly Asn Val Leu Ile Pro Gln Asp Ile Asn Ala Pro Pro Pro Arg Leu Cys Leu Gly Ile Ser Val Asn Glu Pro Asn Asn Lys Asp Asp Gly Lys Ser His Asn His Ser Lys Tyr Ser Asp His Glu Ile Ala Leu Ala Glu Asp Ala Gln Arg Arg Ala Ile Ile Glu Ala Glu Lys His Ala Gln Glu Ala His Arg Gln Ala Lys Lys Ile Ala Gln Glu Val Ser Gly Tyr Arg Ser His Arg Phe Ile Ser Ile Lys Glu Gly Gly Ser Val Gly Asn Ser Asn Thr Asn Gly Asn Asp Asn Asp Glu Asp Asp Asp Glu Val Glu Glu Ala Val Asp Lys Lys Leu Ala Asn Asp Val Ser Val Asp Gly Pro Leu His Val His Glu Gln His Phe Glu Glu Glu Ile Glu Ser Lys Thr Gly Glu Lys Thr Ile Ser Leu Glu Thr

EP99870141.1

Ile Tyr Thr Arg Cys Cys His Leu Arg Glu Ile Leu Pro Ile Pro Ala 

Thr Leu Lys Gln Leu Lys Asn Lys Thr Ala Pro Leu Glu Val Leu Lys 

Met Leu Asn Pro ys Pro Thr Leu Ile Asp Val Leu Ser Phe Ser Asp Phe Ile Ala Ile Thr Pro Ile Asn Thr Val Ile Phe Asp Asn Val Thr Met Thr Thr Glu Met Leu Lys Asn Phe Leu Gly Ser Leu Thr Tyr Asn Lys Gln Leu Glu Lys Leu Ser Leu Arg Asn Val Ser Ile Asp Glu Leu Gly Trp Lys Tyr Leu Cys Glu Phe Leu Ala Thr Asn Lys Thr Val Lys Lys Leu Asp Ile Ser Gln Gln Arg Ile Lys Pro Asp Thr Pro Asp Thr Ser Ile Arg Gly Asn Met Asn Trp Asp Leu Phe Ile Arg Ser Leu Ile Leu Arg Gly Gly Ile Glu Glu Leu Val Ile Asn Gly Cys Lys Leu Ser Asp Ala Ile Phe Glu Lys Phe Ile Asn Gln Ala Val Lys Lys Ser Thr Tyr Arg Leu Gly Ile Ala Gly Ile Asp Leu Asn Val Lys Lys Ser Glu Met Val Thr Ser Trp Leu Thr Asp Gly Asn Ser Gln Cys Val Gly Val Asp Ile Ala Phe Asn Asp Leu Ser Lys Gly Gln Leu Arg Pro Phe Ile Asn Ala Phe Asn Thr Gly Lys Val Asn Asn Leu Val Phe Phe Ser Leu Asn Ser Thr Asn Leu Ser Asn Ile Glu Glu Thr Ser Asp Leu Ile Lys Ser Leu Ile Asn Val Lys Thr Leu Arg Phe Leu Asp Leu Ser Ser Ile 

Pro Asn Ile Phe Pro Lys Ile Ile Thr His Leu Asp Lys Tyr Leu Pro

Arg	Tyr	Pro	Asn	Leu	Arg	Arg	Ile	His	Phe	Asp	Leu	Asn	Glu	Leu	Thr
				885					890					895	

- Ala Gln Ala Ile Gly Ser Leu Ala Gly Cys Leu Ser Lys Met Pro Gln 900 905 910
- Leu Val His Val Ser Leu Leu Gly Asn Arg Asn Leu Ser Thr Thr Ser 915 920 925
- Ala Ala Thr Leu Tyr Gly Ala Val Lys Gln Ser Lys Thr Leu Phe Ala 930 935 940
- Leu Asp Leu Asp Tyr Asp Leu Ile Pro Asp Gln Leu Ser Gln Arg Ile 945 950 955 960
- Ala Phe Tyr Leu Met Arg Asn Leu Glu Tyr Thr Leu Lys Pro Ser His
  965 970 975
- Gly Gly Asn Ile Glu Ser Asn Pro Glu Lys Pro Glu Asp Leu Met Tyr 980 985 990
- Asp Gly Ser Leu Leu Met Glu Thr Ala Glu Lys Leu Leu Val Glu Ile 995 1000 1005
- Glu Lys Gly Lys Lys Glu Asp Ile Lys Met Gln Arg Ile Ile Ser Asp 1010 1015 1020
- Ser Val Leu Glu Arg Thr Arg Ser Ile Arg Lys Asp Ile His Lys Thr 1025 1030 1035 1040
- Ile Asp Thr Leu Phe Glu Gln Arg Asn Leu Gly Lys Leu Ser Phe Glu
  1045 1050 1055
- Gly Lys Glu Asn Leu Val Arg Phe Cys Leu Leu Asp Ser Ser Leu Glu 1060 1065 1070
- Lys Leu Val Val Met Val Glu Glu His Ala Asn Gly Leu Leu Thr 1075 1080 1085
- Pro Thr Thr Ser Thr Asp Asp Leu Arg Ser Arg Ala Met Ser Pro Ser 1090 1095 1100
- Val Thr Val Asp Thr Ile His Glu Ser Ala Asn Glu Leu Ile Thr Ala 1105 1110 1115 1120
- Gly Pro Ile Leu Ser Pro His Val Asn Arg Lys Ala Glu Gln Ser Ser 1125 1130 1135

Tyr Phe Pro Val The Ala Asn Asn Asn Asn Leu Thr Pro His Gln Val 1140 1145 1150

Val Val Glu Ser Asn Asp Glu Gly Arg Asp Val Pro Ile Asp Lys M t 1155 1160 1165

Thr Gly Arg Pro Val Leu Ile Arg Ser Ile Ser Gln Thr Ser Val His 1170 1175 1180

Ala Lys Glu Gln Glu Ile Glu Glu Glu Glu Leu His Lys Phe Gly Phe 1185 1190 1195 1200

Phe Ile Gln Gln Lys Glu Arg Gln Lys Gln Gln Gln Gln Gln Gln Gln 1205 1210 1215

Gln Gln Asn Ser His His Gln His Gln Pro Ala Gln Ser Ile Gln Gln 1220 1225 1230

Glu Asn Gln Ser Pro Ser Pro Gln Gln Gly Lys Tyr Glu Asp Leu Pro 1235 1240 1245

Ile Leu Asn Thr Leu Pro Ser Gly Pro Glu Leu Arg Asp Ala Ile Met 1250 1255 1260

Ala Ala Lys Gly Val Ala Asn Val Thr Glu Leu Ile Asp Arg Ile Asn 1265 1270 1275 1280

Asn His Arg Val Lys Ile Asp Ala Pro Ser Thr Lys His His Glu 1285 1290 1295

Leu Asn Lys Pro Asn Ser Asp Lys Val Val Glu Asp Glu Val Glu Val
1300 1305 1310

Ser Asp Asn Ala Ser Ile Asp Ser Thr Asn Gly Asp Asp Leu His Gln 1315 1320 1325

Leu Gly Asp Gly Lys His Asn Gly Asn Gly Thr Val Asp Pro Met Val 1330 1335 1340

Ser Glu Val Tyr Asp Lys Leu Leu Asn Asp Ala Glu Arg Val Arg Ser 1345 1350 1355 1360

Asn Arg Asp Ile

<210> 453

<211> 1859

```
<212> DNA
<213> Candida albicans
<400> 453
acggaataat gttatgcgaa cagagtaact gggattatca acaagcatca gtcaatttca 60
aaaactcggc tgcttctttg cctagtgatg catttgtaca atagaattat aattaaaaga 120
aaggtagaat agaattcaat gtaggtgata tcactttctg agagttctta ttacgatata 180
taaaatacat agccaaaatt ataatgaagt aaaaacatgt aactgtgtaa ttttattcaa 240
gtccaaaggt attgattaat attgtagagt gttgagcatt taaaatatga aggaagaccg 300
ataacctaaa gttttcctca gggtgtcgaa ggctaggagg aataaaatct gagtagagaa 360
aacgaattac acaagcacat cctaaacacc actctgttgg agcaccaatc aacctgagaa 480
atgcaaactt gtaataaaat atgggataca atatagcaat ggtaacagat tttttttacc 540
ctcaacctgg aggagtagag tttcatgtgt atcatttatc acaaaaactc attgaactag 600
gacactcagt ggttatcata actcataatt attcatcaag aaatggtgta cgagtattaa 660
cgaatggttt gaaagtgtat tatgtaccac tttgggtgat ctatagaagc tcagttttcc 720
caactgtatt tctgtgcttc ccaatattga ggaatatctt catacgagaa aacattgaga 780
ttattcacgg acatggttcc ttcagcacat tatgccacga agctatatta catggccgaa 840
caatgggatt aaaaacagtc ttcactgatc attcactttt tggatttgcc gagattggat 900
caattatggg gaataaagca ttaaagttca ctttcagtga tgttggccat gttatctgtg 960
tcagtcacac ctgtaaagaa aacacggttt taagaggatc aatagacccc ataaaagtga 1020
gtgtgatacc gaatgcagtt attctgaaag atttcaagcc caaatcgcat tgtgttaaca 1080
agaactatac taaagagatc accattgtgg tgatcacgag attgtttcca aataaaggag 1140
ccgatctatt aacggctgtt atccccaaaa tttgccagtt gaaaccaaaa gtgaaatttc 1200
taattgctgg tgacggcccc aagtttttag atttagaaca aatgagagaa aagtactttc 1260
ttcaggaaag ggttacatta gtaggcgcta taaaacacga agaagtaaga gatgtaatgg 1320
tccaaggtga catatactta catccttcat taacagaggc gtttggtaca gttattgtgg 1380
aagctgcatc atgtgggtta tatgttgtca ctacaaaagt tggaggcata cccgaagtct 1440
taccaaacqa aatgacaagc tttgctgaac cggaagaaaa ctcacttatt gatgctgcta 1500
tagatgctat aaataaaatt gaaagtaatg aaatcgatac ctcaaaattt cacgatgcgg 1560
ttgcaaagat gtacagttgg aatgatattg caagaagaac agaaaatgtt tataattcac 1620
ttgatttaga caaactaaac gagtctttac ttcaccgatt acaaagatac tattgttgtg 1680
gtataatagc aggcaaactt tatgctttat gtgtaatagt ggatattttt attttcgtga 1740
tactagaatg gttgtatccc gctgatcata tcgataaagc aacaaaatgg ccactggcta 1800
tcaaggaaga agacgagctg gaagaagaaa catttatttt tccgaacaaa gtaaattag 1859
<210> 454
<211> 452
<212> PRT
<213> Candida albicans
<400> 454
```

Gly Gly Val Glu Phe His Val Tyr His Leu Ser Gln Lys Leu Ile Glu

Met Gly Tyr Asn Ile Ala Met Val Thr Asp Phe Phe Tyr Pro Gln Pro

10

5

550

			20	,				23				· Care	, 30		
Let	ı Gly	His		Val	Val	Ile	Ile 40	Thr	His	Asn	Tyr	Ser 45		Arg	Asn
Gly	7 Val 50		Val	Leu	Thr	Asn 55	Gly	Leu	Lys	Val	Tyr 60	Tyr	Val	Pro	Leu
Trp 65	Val	Ile	Tyr	Arg	Ser 70	Ser	Val	Phe	Pro	Thr 75	Val	Phe	Ser	Cys	Phe 80
Pro	Ile	Leu	Ara	Asn	_Ile	Phe	Ile	Ara	Glu	Asn	Ile	Glu	Ile	Ile	His
				85					90					95	
Gly	His	Gly	Ser 100		Ser	Thr	Leu	Cys 105	His	Glu	Ala	Ile	Leu 110	His	Gly
Arg	Thr	Met 115	Gly	Leu	Lys	Thr	Val 120	Phe	Thr	Asp	His	Ser 125	Leu	Phe	Gly
Ph∈	Ala 130	Glu	Ile	Gly	Ser	Ile 135	Met	Gly	Asn	Lys	Ala 140	Leu	Lys	Phe	Thr
Phe 145	Ser	Asp	Val	Gly	His 150	Val	Ile	Суѕ	Va1	Ser 155	His	Thr	Суѕ	Lys	Glu 160
Asn	Thr	Val	Leu	Arg 165	Gly	Ser	Ile	Asp	Pro 170	Ile	Lys	Val	Ser	Val 175	Ile
Pro	Asn	Ala	Val 180	Ile	Ser	Lys	Asp	Phe 185	Lys	Pro	Lys	Ser	His 190	Cys	Val
Asn	Lys	Asn 195	Tyr	Thr	Lys	Glu	Ile 200	Thr	Ile	Val	Val	11e 205	Thr	Arg	Leu
Phe	Pro 210	Asn	Lys	Gly	Ala	Asp 215	Leu	Leu	Thr	Ala	Val 220	Ile	Pro	Lys	Ile
Cys 225	Gln	Leu	Lys	Pro	Lys 230	Va1	Lys	Phe	Leu	11e 235	Ala	Gly	Asp	Gly	Pro 240
Lys	Phe	Leu	Asp	Leu 245	Glu	Gln	Met	Arg	Glu 250	Lys	Tyr	Phe	Leu	Gln 255	Glu
Arg	Val	Thr	Leu	Val	Gly	Ala	Ile	Lys	His	Glu	Glu	Val	Arg	Asp	Val

Met Val Gln Gly Asp Ile Tyr Leu His Pro Ser Leu Thr Glu Ala Phe

# EP99870141.1

SEQL / "

275

285

Gly Thr Val Ile Val Glu Ala Ala Ser Cys Gly Leu Tyr Val Val Thr 290 295 300

280

Thr Lys Val Gly Gly Ile Pro Glu Val Leu Pro Asn Glu Met Thr Ser 305 310 315 320

Phe Ala Glu Pro Glu Glu Asn Ser Leu Ile Asp Ala Ala Ile Asp Ala 325 330 335

Ile Asn Lys Ile Glu Ser Asn Glu Ile Asp Thr Ser Lys Phe His Asp 340 345 350

Ala Val Ala Lys Met Tyr Ser Trp Asn Asp Ile Ala Arg Arg Thr Glu 355 360 365

Asn Val Tyr Asn Ser Leu Asp Leu Asp Lys Leu Asn Glu Ser Leu Leu 370 375 380

His Arg Leu Gln Arg Tyr Tyr Cys Cys Gly Ile Ile Ala Gly Lys Leu 385 390 395 400

Tyr Ala Leu Cys Val Ile Val Asp Ile Phe Ile Phe Val Ile Leu Glu
405 410 415

Trp Leu Tyr Pro Ala Asp His Ile Asp Lys Ala Thr Lys Trp Pro Ser 420 425 430

Ala Ile Lys Glu Glu Asp Glu Ser Glu Glu Glu Thr Phe Ile Phe Pro 435 440 445

Asn Lys Val Asn 450

<210> 455

<211> 1231

<212> DNA

<213> Candida albicans

<400> 455

gatacaattc acgacctcta atttccttag tgataggtta atcatgaaat atataaaatg 60 tctagaaata attgaataat gatataaata taatgttgtg aaacttgtgg aaaaatttga 120 cagcctagta actcaagtgt tgttacactc ttgttattat tattattata tcgtttacaa 180 gtagatttct cattttgaac agcaaatact gtcgttaata ggaatcagag gcagaaagaa 240 agagagagaa aaaaaaaaag gacacattta cacgtacacc cttaacttga aggaaaaaaa 300

caaaacaaga gaalaaga gaaaatactt tcaaca a aagattgaga 360 tattggtgat tttcaaca c caaaaaaaaa gaatacacaa cttgta 2t aacagaattt 420 gaacatttct agagtttctt atattccccc caaggtggac aattaataat aacatttgat 480 tatacaagaa catttcaatc atgtggattt ttgactggtg tatgtaatga tttatcttta 540 ccgaatcaat actttatta tgagtgattg gttggttggt tataatacca ccactttatt 600 ctattaacca agtttgatct aattactgat ctgtatatac taaccaaata tttaccttat 660 cacttaatta tttacagttc aagatatatt atcatcatta ggattatgga ataaacatgc 720 caaattatta tttttagggt tagataatgc tggtaaaact actcttttac atatgttaaa 780 gaatgataga ttggccactt tacaaccaac attacatcca acttcagaag aattggccat 840 tggatcagtt agatttacta cttttgattt aggtggacat caacaagcta gaagattatg 900 gaaagattat ttccctgaag tcaatggtat tgtcttttta gtcgatgctg ctgataccga 960 aagatttgct gaatccaaag ctgaattgga aagtttattt agaattgaag aattgagtca 1020 agttccattt gttattttgg gtaataagat tgatgttcct actgcagtag gggaaatgga 1080 attgaaaaat gcccttggat tatataatac tactggtaaa gatactggta aattgcctga 1140 aggtactaga ccaattgaag tgtttatggt ttccgttgtt atgagatctg gatatggtga 1200 agccttcaaa tggttatcac aatacattta a 1231

<210> 456

<211> 190

<212> PRT

<213> Candida albicans

<400> 456

Met Trp Ile Phe Asp Trp Val Gln Asp Ile Leu Ser Ser Leu Gly Leu 1 5 10 15

Trp Asn Lys His Ala Lys Leu Leu Phe Leu Gly Leu Asp Asn Ala Gly
20 25 30

Lys Thr Thr Leu Leu His Met Leu Lys Asn Asp Arg Leu Ala Thr Leu 35 40 45

Gln Pro Thr Leu His Pro Thr Ser Glu Glu Leu Ala Ile Gly Ser Val
50 55 60

Arg Phe Thr Thr Phe Asp Leu Gly Gly His Gln Gln Ala Arg Arg Leu 65 70 75 80

Trp Lys Asp Tyr Phe Pro Glu Val Asn Gly Ile Val Phe Leu Val Asp 85 90 95

Ala Ala Asp Thr Glu Arg Phe Ala Glu Ser Lys Ala Glu Leu Glu Ser 100 105 110

Leu Phe Arg Ile Glu Glu Leu Ser Gln Val Pro Phe Val Ile Leu Gly 115 120 125

## EP99870141.1





Asn Lys Ile Asp Val Pro Thr Ala Val Gly Glu Met Glu Leu Lys Asn 130 135 140

Ala Leu Gly Leu Tyr Asn Thr Thr Gly Lys Asp Thr Gly Lys Leu Pro 145 150 155 160

Glu Gly Thr Arg Pro Ile Glu Val Phe Met Val Ser Val Val Met Arg 165 170 175

Ser Gly Tyr Gly Glu Ala Phe Lys Trp Leu Ser Gln Tyr Ile 180 185 190